SPARSITY EXPLOITATION IN THE EXTENDED NAPHTALI-SANDHOLM METHOD FOR SOLUTION OF INTERLINKED MULTISTAGED SEPARATORS

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by
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to the

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CERTIFICATE

It is certified that the present work entitled,
"SPARSITY EXPLOITATION IN THE EXTENDED NAPHTALI—SANDHOLM
METHOD FOR SOLUTION OF INTERLINKED MULTISTAGED SEPARATORS"
has been carried out by Mr. Manoj Kumar Jain under my
supervision and that it has not been submitted elsewhere for
a degree.

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-Manoj Kumar Jain

ABSTRACT

An efficient algorithm has been presented for solving the multicomponent, multistage separation process problems involving single columns, columns with pump-around or bypass, or a system of interlinked columns. In this algorithm which is an extension of the Naphtali-Sandholm method, the sparsity and the structure of the submatrices in the Jacobian are exploited, while performing the matrix multiplications and inversions. The operation count was performed for the various matrix multiplications and inversions involved in the algorithm, and using a variety of test-problems it has been shown that the sparsity exploitation results in a significant reduction in the computational and storage requirements. The saving in the computations improves with the increase in the number of components. An efficient approach has been proposed to solve problems with intermediate tray specifications, in which the tridiagonal band structure of the Jacobian is preserved, and has been shown to be more advantageous in both the computational and storage aspects, than the one proposed by Hofeling and Seader:

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CHAPTER 1

INTRODUCTION

The separation processes like distillation, absorption and extraction etc. are widely used in the chemical industries to fractionate the multicomponent mixtures. Complex systems of interlinked columns have been shown to frequently achieve more economic and effective separation of multicomponent mixtures than the conventional single columns and their sequential arrangements. In view of the increasing energy and raw material cost, a continuous evaluation of the performance is required for economic operation as well as to maintain the quality of the product. This requires simulation of multicomponent interlinked columns.

The Naphtali-Sandholm method [1] is widely used for the simulation of single columns due to its good convergence characteristics. For the simulation of interlinked columns a simultaneous rather than a sequential approach is preferred. That is, rather than repeatedly solving a sequence of single column problems until the solution to the entire system is obtained, it is preferred to solve for all the columns simultaneously.

In one version of this approach, the methods like capital—theta method of Holland [10], the equations describing each column are considered separately but all these single

column problems are converged simultaneously rather than repeatedly one at a time. Whereas the more powerful approach is to consider the equations describing all the separators simultaneously, and solve by the Newton-Raphson or a similar method with good convergence characteristics. Following this approach several methods have been developed by Hofeling and Seader [2], Kubicek [7,12], Stadtherr [5], Browne [13] etc. While these methods differ in their formulation of equations and their selection of independent variables, all rely on reordering some or all of the linearized equations to produce an almost-band or almost-block-tridiagonal coefficient matrix. Whereas these methods differ in their approach to obtain the desired matrix form and their method of solving the reordered linear system.

The Naphtali-Sandholm method which was originally formulated to solve the single column problems has been extended [2], to handle the system of interlinked columns, and the columns with pump-around or bypass, while retaining the technique of total linearization and simultaneous solution of all the equations in the system by the Newton-Raphson method.

Though this method offers a good convergence characteristics, the large computational and storage requirements are
the impediments.

When the Newton-Raphson technique is applied to solve

the equations in the Naphtali-Sandholm method, the resulting Jacobian matrix has a tridiagonal-band or almost-tridiagonal-band structure with a few off-tridiagonal elements. The submatrices in the Jacobian are sparse and have a definite structure.

It appears from the literature that the sparsity of the submatrices has not been exploited. The objective of this work is to present an efficient algorithm, which takes the advantage of the sparsity and the structure of the submatrices in the Jacobian, to solve the separation process problems.

In Chapter 2, first the method of formulation and solution for single column problems has been presented and then extended to the case of interlinked columns. Chapter 3 deals with the exploitation of sparsity in the various matrix multiplications and inversions. In Chapter 4, an efficient approach has been proposed to solve the problems with the intermediate tray specifications, and a variety of specifications for condensers, reboilers and intermediate trays have been presented. Chapter 5 deals with the details of implementation of separation process problems on the computer. The results of various test-problems and a brief discussion have been presented in Chapter 6 and the conclusions in Chapter 7.

CHAPTER 2

MODELLING OF MULTICOMPONENT MULTISTAGE COLUMNS

In this chapter, first the modelling of a single multicomponent multistage separation column is presented, which is later extended to a system of interlinked columns. In both the cases, the model proposed by Naphtali and Sandholm [1], is followed, except for a minor reordering of variables and discrepancy function in the formulation of the Jacobian-matrix, the reason for which it is done is explained later.

2.1 Generalized Tray Model for a Single Column

A model for a general tray j, with streams to and from the neighbouring trays j-1 and j+1, and with side streams is shown in Figure 1. The discrepancy functions formulated using mass balances, equilibrium relations and energy balance are as under:

Component Mass Balances:

$$M_{j,i} = l_{j-1,i} + v_{j+1,i} - (1+s_j)l_{j,i} - (1+s_j)v_{j,i}+f_{j,i}$$

for $1 \le i \le C$ and $1 \le j \le N$

(2.1)

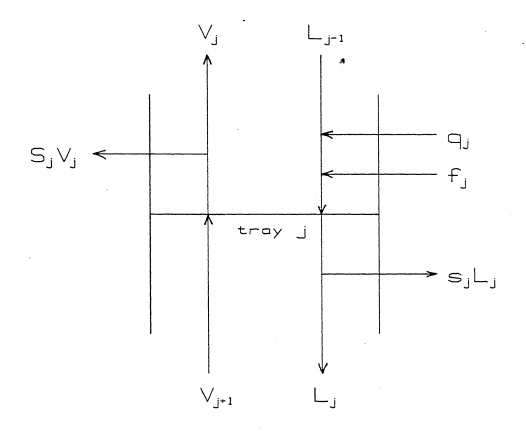


FIG 1 A GENERAL TRAY IN A SINGLE COLUMN

Equilibrium Relationships:

$$Q_{j,i} = \frac{\eta_{j,i}}{L_{j}} l_{j,i} - \frac{v_{j,i}}{V_{j}} + \frac{(1 - \eta_{j})}{V_{j+1}} v_{j+1,i}$$

$$(2.2)$$
for $1 \le i \le C$ and $1 \le j \le N$

where

$$\eta_{j} = \frac{Y_{j,i} - Y_{j+1,i}}{K_{j,i} X_{j,i} - Y_{j+1,i}}$$
 (2.3)

Energy Balance:

$$E_{j} = \sum_{i=1}^{c} h_{j-1,i} l_{j-1,i} + \sum_{i=1}^{c} H_{j+1,i} v_{j+1,i} - (1+s_{j})$$

$$\sum_{i=1}^{c} h_{j,i} l_{j,i}$$

$$- (1+s_{j}) \sum_{i=1}^{c} H_{j,i} v_{j,i} + \sum_{i=1}^{c} h_{F_{j,i}} f_{j,i} + q_{j}$$

$$for 1 < j < N$$

$$(2.4)$$

where, $l_{j,i}$ and $v_{j,i}$ are the molar liquid and vapor flow rates leaving the tray j after the liquid and vapor side streams $s_j l_{j,i}$ and $s_j v_{j,i}$ have been drawn from stage j.

Thus there are (2c+1) equations (discrepancy functions), viz. $M_{j,i}$, $Q_{j,i}$ and E_{j} , and (2c+1) variables, viz. $l_{j,i}$, $v_{j,i}$ and T_{j} for each tray and hence a total of N(2c+1) of each for the whole column. The set of these N(2c+1) discrepancy functions and variables may be written in a compact form as

follows:

$$\vec{F}(\vec{X}) = \vec{0} \tag{2.5}$$

where,

$$\vec{F} = (\vec{F}_1, \vec{F}_2, \dots, \vec{F}_N)^T \qquad (2.6)$$

$$\vec{F}_{j} = (M_{j,1}, M_{j,2}, ..., M_{j,c}, Q_{j,1}, ..., Q_{j,c}, E_{j})^{T}$$

for 1 < j < N

and

$$\overline{X} = (\overline{x}_1, \overline{x}_2, \dots, \overline{x}_j, \dots, \overline{x}_N)^T$$
 (2.8)

$$\bar{x}_{j} = (l_{j,1}, l_{j,2}, ..., l_{j,c}, v_{j,1}, v_{j,2}, ..., v_{j,c}, T_{j})^{T}$$

(2.9)

for $1 \le j \le N$

It may be noted here that:

- (a) The ordering of the discrepancy functions and variables is a bit different from the one proposed by Naphtali and Sandholm [1]. This reordering results in the suitable structure of submatrices for the purpose of sparsity exploitation, which is explained later.
- (b) A minor change has been made in the discrepancy function $Q_{j,i}$, which was proposed by Naphatali and Sandholm as: $Q_{j,i} = \frac{\eta_j K_{j,i} V_j}{L_i} 1_{j,i} V_{j,i} + \frac{(1 \eta_j)V_j}{V_{j+1}} V_{j+1,i}$

(2.10)

The discrepancy function (2.2) is obtained by dividing the original equation (2.10) by V_j . This minor change has improved the convergence characteristics.

Employing the Newton-Raphson method to solve the system of equations (2.5) simultaneously we obtain:

$$\Delta \overline{x}_{m+1} = -\left(\frac{d\overline{F}}{d\overline{X}}\right)^{-1} \overline{F}_{m} \qquad (2.11)$$

$$= - \overline{J}_{m}^{-1} \overline{F}_{m} \qquad (2.12)$$

where, \overline{J} is the Jacobian matrix and m is the iteration number, and,

$$\overline{x}_{m+1} = \overline{x}_m + \Delta \overline{x}_{m+1} \qquad (2.13)$$

The structure of the Jacobian matrix is:

$$\begin{bmatrix} \overline{\mathbb{B}}_1 & \overline{\mathbb{C}}_1 \\ \overline{\mathbb{A}}_2 & \overline{\mathbb{B}}_2 & \overline{\mathbb{C}}_2 \\ \overline{\mathbb{A}}_3 & \overline{\mathbb{B}}_3 & \overline{\mathbb{C}}_3 \\ & & & & & & \\ & & & \overline{\mathbb{A}}_j & \overline{\mathbb{B}}_j & \overline{\mathbb{C}}_j \\ & & & & \overline{\mathbb{A}}_N & \overline{\mathbb{B}}_N \end{bmatrix}$$

where,

$$\overline{\overline{A}}_{j} = \frac{\partial \overline{F}_{j}}{\partial \overline{x}_{j-1}}, \overline{\overline{B}}_{j} = \frac{\partial \overline{F}_{j}}{\partial \overline{x}_{j}}, \overline{\overline{C}}_{j} = \frac{\partial \overline{F}_{j}}{\partial \overline{x}_{j+1}}$$
 (2.15)

The submatrices $\bar{\bar{A}}_j$, $\bar{\bar{B}}_j$ and $\bar{\bar{C}}_j$ have the following elements:

$$\bar{A}_{j} = \begin{bmatrix}
\frac{\partial M_{j,1}}{\partial 1_{j-1,1}} & \frac{\partial M_{j,1}}{\partial 1_{j-1,c}} & \frac{\partial M_{j,1}}{\partial v_{j-1,1}} & \frac{\partial M_{j,1}}{\partial v_{j-1,c}} & \frac{\partial M_{j,1}}{\partial 1_{j-1}} \\
\frac{\partial M_{j,c}}{\partial 1_{j-1,1}} & \frac{\partial M_{j,c}}{\partial 1_{j-1,c}} & \frac{\partial M_{j,c}}{\partial v_{j-1,1}} & \frac{\partial M_{j,c}}{\partial v_{j-1,c}} & \frac{\partial M_{j,c}}{\partial 1_{j-1}} \\
\frac{Q_{j,1}}{1_{j-1,1}} & \frac{\partial Q_{j,1}}{\partial 1_{j-1,c}} & \frac{\partial Q_{j,1}}{\partial v_{j-1,1}} & \frac{\partial Q_{j,1}}{\partial v_{j-1,c}} & \frac{\partial Q_{j,1}}{\partial 1_{j-1}} \\
\frac{\partial Q_{j,c}}{\partial 1_{j-1,1}} & \frac{\partial Q_{j,c}}{\partial 1_{j-1,c}} & \frac{\partial Q_{j,c}}{\partial v_{j-1,1}} & \frac{\partial Q_{j,c}}{\partial v_{j-1,c}} & \frac{\partial Q_{j,c}}{\partial 1_{j-1}} \\
\frac{\partial E_{j}}{\partial 1_{j-1,1}} & \frac{\partial E_{j}}{\partial 1_{j-1,c}} & \frac{\partial E_{j}}{\partial v_{j-1,1}} & \frac{\partial E_{j}}{\partial v_{j-1,c}} & \frac{\partial E_{j}}{\partial 1_{j-1}} \\
\frac{\partial E_{j}}{\partial 1_{j-1,1}} & \frac{\partial E_{j}}{\partial 1_{j-1,c}} & \frac{\partial E_{j}}{\partial v_{j-1,1}} & \frac{\partial E_{j}}{\partial v_{j-1,c}} & \frac{\partial E_{j}}{\partial 1_{j-1,c}} \\
\frac{\partial E_{j}}{\partial 1_{j-1,c}} & \frac{\partial E_{j}}{\partial 1_{j-1,c}} & \frac{\partial E_{j}}{\partial v_{j-1,1}} & \frac{\partial E_{j}}{\partial v_{j-1,c}} & \frac{\partial E_{j}}{\partial 1_{j-1,c}} \\
\frac{\partial E_{j}}{\partial 1_{j-1,c}} & \frac{$$

$$\bar{B}_{j} = \begin{bmatrix} \frac{\partial M_{j,1}}{\partial I_{j,1}} & \frac{\partial M_{j,1}}{\partial I_{j,1}} & \frac{\partial M_{j,1}}{\partial V_{j,1}} & \frac{\partial M_{j,1}}{\partial V_{j,c}} & \frac{\partial M_{j,1}}{\partial I_{j}} \\ \frac{\partial M_{j,c}}{\partial I_{j,1}} & \frac{\partial M_{j,c}}{\partial I_{j,c}} & \frac{\partial M_{j,c}}{\partial V_{j,1}} & \frac{\partial M_{j,c}}{\partial V_{j,c}} & \frac{\partial M_{j,c}}{\partial I_{j}} \\ \frac{\partial Q_{j,1}}{\partial I_{j,1}} & \frac{\partial Q_{j,1}}{\partial I_{j,c}} & \frac{\partial Q_{j,1}}{\partial V_{j,1}} & \frac{\partial Q_{j,1}}{\partial V_{j,c}} & \frac{\partial Q_{j,c}}{\partial I_{j}} \\ \frac{\partial Q_{j,c}}{\partial I_{j,1}} & \frac{\partial Q_{j,c}}{\partial I_{j,c}} & \frac{\partial Q_{j,c}}{\partial V_{j,1}} & \frac{\partial Q_{j,c}}{\partial V_{j,c}} & \frac{\partial Q_{j,c}}{\partial I_{j}} \\ \frac{\partial E_{j}}{\partial I_{j,1}} & \frac{\partial E_{j}}{\partial I_{j,c}} & \frac{\partial E_{j}}{\partial V_{j,1}} & \frac{\partial M_{j,1}}{\partial V_{j,c}} & \frac{\partial M_{j,1}}{\partial V_{j,c}} & \frac{\partial M_{j,1}}{\partial I_{j}} \\ \frac{\partial M_{j,c}}{\partial I_{j+1,1}} & \frac{\partial M_{j,c}}{\partial I_{j+1,c}} & \frac{\partial M_{j,c}}{\partial V_{j+1,1}} & \frac{\partial M_{j,1}}{\partial V_{j+1,c}} & \frac{\partial M_{j,1}}{\partial V_{j+1,c}} & \frac{\partial M_{j,1}}{\partial V_{j+1,c}} & \frac{\partial M_{j,1}}{\partial V_{j+1,c}} \\ \frac{\partial Q_{j,c}}{\partial I_{j+1,1}} & \frac{\partial Q_{j,1}}{\partial I_{j+1,c}} & \frac{\partial Q_{j,1}}{\partial V_{j+1,1}} & \frac{\partial Q_{j,1}}{\partial V_{j+1,c}} & \frac{\partial Q_{j,c}}{\partial I_{j+1,c}} \\ \frac{\partial Q_{j,c}}{\partial I_{j+1,c}} & \frac{\partial Q_{j,c}}{\partial V_{j+1,1}} & \frac{\partial Q_{j,c}}{\partial V_{j+1,c}} & \frac{\partial Q_{j,c}}{\partial I_{j+1,c}} \\ \frac{\partial E_{j}}{\partial I_{j+1,1}} & \frac{\partial E_{j}}{\partial I_{j+1,c}} & \frac{\partial E_{j}}{\partial V_{j+1,1}} & \frac{\partial E_{j}}{\partial V_{j+1,c}} & \frac{\partial Q_{j,c}}{\partial I_{j+1,c}} \\ \frac{\partial E_{j}}{\partial I_{j+1,1}} & \frac{\partial E_{j}}{\partial I_{j+1,c}} & \frac{\partial E_{j}}{\partial V_{j+1,1}} & \frac{\partial E_{j}}{\partial V_{j+1,c}} & \frac{\partial E_{j}}{\partial I_{j+1,c}} \\ \frac{\partial E_{j}}{\partial I_{j+1,c}} & \frac{\partial E_{j}}{\partial V_{j+1,1}} & \frac{\partial E_{j}}{\partial V_{j+1,1}} & \frac{\partial E_{j}}{\partial V_{j+1,c}} & \frac{\partial E_{j}}{\partial I_{j+1,c}} \\ \frac{\partial E_{j}}{\partial I_{j+1,c}} & \frac{\partial E_{j}}{\partial I_{j+1,c}} & \frac{\partial E_{j}}{\partial V_{j+1,1}} & \frac{\partial E_{j}}{\partial V_{j+1,c}} & \frac{\partial E_{j}}{\partial I_{j+1,c}} \\ \frac{\partial E_{j}}{\partial I_{j+1,c}} & \frac{\partial E_{j}}{\partial I_{j+1,c}} & \frac{\partial E_{j}}{\partial V_{j+1,1}} & \frac{\partial E_{j}}{\partial V_{j+1,c}} & \frac{\partial E_{j}}{\partial V_{j+1,c}} \\ \frac{\partial E_{j}}{\partial I_{j+1,c}} & \frac{\partial E_{j}}{\partial I_{j+1,c}} \\ \frac{\partial E_{j}}{$$

In compact form, the structures of $\overline{\overline{A}}$, $\overline{\overline{\overline{B}}}$ and $\overline{\overline{\overline{C}}}$ submatrices may be written as

$$\bar{A}_{j} = \begin{bmatrix}
\bar{T}_{c} & \bar{o}_{c} & \bar{o}_{c} \\
\bar{o}_{c} & \bar{o}_{c} & \bar{o}_{c} \\
\bar{X}_{c} & \bar{o}_{c} & X
\end{bmatrix} (2.19)$$

$$\bar{B}_{j} = \begin{bmatrix}
-(1+s_{j})\bar{T}_{c} & -(1+s_{j})\bar{T}_{c} & \bar{o}_{c} \\
\bar{X}_{c} & \bar{X}_{c} & \bar{X}_{c} \\
\bar{X}_{c} & \bar{X}_{c} & X
\end{bmatrix} (2.20)$$

$$\bar{B}_{j} = \begin{bmatrix}
\bar{o}_{c} & \bar{T}_{c} & \bar{o}_{c} \\
\bar{o}_{c} & \bar{X}_{c} & \bar{o}_{c} \\
\bar{o}_{c} & \bar{X}_{c} & X
\end{bmatrix} (2.21)$$

$$\overline{\overline{C}}_{j} = \begin{bmatrix} \overline{\overline{C}}_{c} & \overline{\overline{T}}_{c} & \overline{\overline{C}}_{c} \\ \overline{\overline{C}}_{c} & \overline{\overline{X}}_{c} & \overline{\overline{C}}_{c} \\ \overline{\overline{C}}_{c} & \overline{\overline{X}}_{c} & x \end{bmatrix}$$
 (2.21)

where.

is an identity matrix of order cxc

is a null matrix of order cxc

is either a row or a column null vector (whose meaning is clear from its position) of order c

- \overline{X}_{c} is a matrix with non-zero elements, of order cxc
- $\overline{X}_{\mathbf{c}}$ is either a row or a column vector with non-zero elements, of order \mathbf{c}
- X is a non-zero element.

It may be noted that the Jacobian matrix in (2.14) has a tridiagonal-band-structure and is highly sparse. This is due to the fact that only those derivatives are non-zero which are obtained by differentiating the discrepancy functions of any tray j, with respect to the variables of the same tray j, or the variables of the neighbouring trays j-1 or j+1.

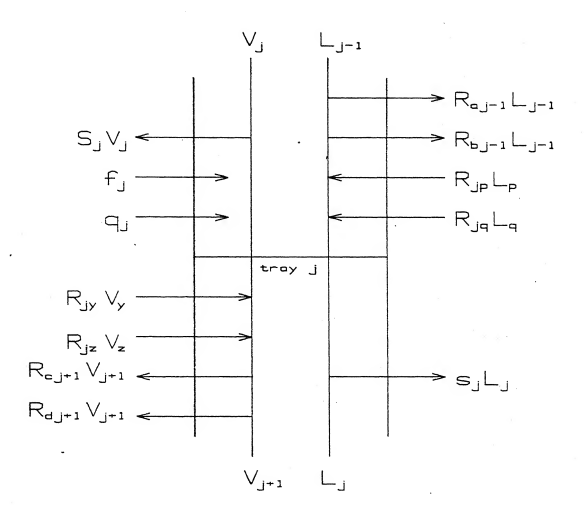
Further, the submatrices \overline{A} and $\overline{\overline{C}}$ in the Jacobian $\overline{\overline{J}}$ are also sparse which may be noted in the structures presented in (2.19) and (2.21).

2.2 Interlinked Columns:

The Naphtali-Sandholm method which was originally formulated to solve the single column problems, can be extended to handle a system of interlinked columns, and the columns with pump-around or bypass.

Generalized Tray Model:

A model for a general tray j, in a system of interlinked columns or in the columns with bypass or pump-around streams is presented in Figure 2. In addition to the streams to and from the neighbouring trays and side-streams, this model



$$\Gamma_{L_{j-1}} = 1 - (R_{a_{j-1}} + R_{b_{j-1}} + \dots)$$

$$\Gamma_{V_{j+1}} = 1 - (R_{a_{j+1}} + R_{a_{j+1}} + \dots)$$

FIG 2 A GENERAL TRAY

IN A SYSTEM OF

INTERLINKED COLUMNS

includes the interlinked-streams which link the tray j to trays which are not its immediate neighbours in the system-model.

Let $R_{jp} L_p$, $R_{jq} L_q$, $R_{jy} V_y$, $R_{jz} V_z$... be the streams which are leaving the tray p, q, y, z ..., and are fed to the stage j, where R_{kj} represents the fraction of stream which is leaving the tray k and fed to tray j, and

- $R_{kj} = 0$ if there is no stream from tray k to tray j
 - = 1 if the whole stream from k is fed to tray j
 - = a a value between 0 and 1 if a part of the stream leaving the tray k is fed to tray j.

The sum of all liquid and vapor interlinked streams leaving the tray j is represented by $(1 - r_{L_{j+1}})L_{j-1}$ and $(1 - r_{V_{j+1}})V_{j+1}$, respectively.

The discrepancy functions formulated based on the component material balances, equilibrium relationships and enthalpy balance are:

Component Mass Balances:

$$M_{j,i} = r_{L_{j-1}} l_{j-1,i} + r_{V_{j+1}} v_{j+1,i} - (1+s_{j}) l_{j,i} - (1+s_{j}) v_{j,i}$$

$$+ R_{jp} l_{p,i} + R_{jq} l_{q,i} + \cdots + R_{jy} v_{y,i} + R_{jz} v_{z,i} + \cdots$$

$$+ f_{j,i} \qquad \text{for } 1 \le i \le C \text{ and } 1 \le j \le N \qquad (2,22)$$

Equilibrium Relationships:

$$Q_{j,i} = \frac{\eta_{j}^{K}_{j,i}}{L_{j}} l_{j,i} - \frac{v_{j,i}}{v_{j}} + \frac{(1 - \eta_{j})r_{V_{j+1}}v_{j+1}v_{j+1,i}}{r_{V_{j+1}}v_{j+1}^{H}} v_{y} v_{y} + R_{jz}v_{z}$$

$$+ \frac{(1 - \eta_{j})R_{jy}v_{y,i}}{r_{V_{j+1}}v_{j+1}^{H}} + R_{j}v_{y}^{V}v_{y}^{H} v_{j}^{H}v_{z}^{U}z}$$

$$+ \frac{(1 - \eta_{j})R_{jz}v_{z,i}}{r_{V_{j+1}}v_{j+1}^{H}} + R_{j}v_{y}^{V}v_{y}^{H} v_{j}^{H}z^{U}z} + \dots (2.23)$$

$$+ \frac{v_{j+1}}{r_{V_{j+1}}} v_{j+1}^{H} + R_{j}v_{y}^{H}v_{y}^{H}v_{j}^{H}z^{U}z} + \dots (2.23)$$

$$+ \frac{v_{j+1}}{r_{V_{j+1}}} v_{j+1}^{H} + R_{j}v_{y}^{H}v_{y}^{H}v_{j}^{H}z^{U}z} + \dots (2.23)$$

Enthalpy Balance:

$$E_{j} = r_{L_{j-1}} \sum_{i=1}^{c} h_{j-1,i} l_{j-1,i} + r_{V_{j+1}} \sum_{i=1}^{c} H_{j+1,i} v_{j+1,i}$$

$$-(1+s_{j}) \sum_{i=1}^{c} h_{j,i} l_{j,i} - (1+s_{j}) \sum_{i=1}^{c} H_{j,i} v_{j,i}$$

$$+ R_{jp} \sum_{i=1}^{c} h_{p,i} l_{p,i} + R_{jq} \sum_{i=1}^{c} h_{q,i} l_{q,i} + \cdots$$

$$+ R_{jy} \sum_{i=1}^{c} H_{y,i} v_{y,i} + R_{jz} \sum_{i=1}^{c} H_{z,i} v_{z,i} + \cdots$$

$$+ \sum_{i=1}^{c} h_{F_{j,i}} f_{j,i} + q_{j} \text{ for } 1 \leq j \leq N \quad (2.24)$$

The choice of variable remains the same as it was for a single column viz. $l_{j,i}$, $v_{j,i}$ and T_{j} , in that order. Thus,

we have N(2c+1) equations and N(2c+1) variables which are to be solved simultaneously using the Newton-Raphson method.

It may be noted here that the discrepancy functions of the tray j may also have the variables of the trays, which are not the immediate neighbours of the tray j, due to the presence of interlinked-streams, bypass or pump-arounds. Therefore, the Jacobian matrix may have some off-tridiagonal blocks in addition to the tridiagonal-band. The position and structure of these off-tridiagonal blocks depend upon the arrangement of the columns in the system model (Hildalgo and Seader [4]), position of the interlinked trays, and the direction and type (liquid or vapor) of the interlinked streams.

The single stream between two trays (not neighbours) may be classified into four basic types:

- (i) liquid stream from an upper tray to a lower tray
- (ii) liquid stream from a lower tray to an upper tray
- (iii) vapor stream from a lower tray to an upper tray
- (iv) vapor stream from an upper tray to a lower tray.

The reciprocal streams (in which a stream from one stage to another stage, is matched by a stream of the other phase flowing in the opposite direction between the same two stages), may be handled as a combination of these four types of streams.

Consider an arrangement of interlinked-columns in which j and q are the two trays such that j is above q and $q-j \ge 2$, and a stream linking the two. The presence of this stream will cause an off-diagonal block to occur in the Jacobian matrix, and the position and structure of which may be determined considering the four types discussed below:

Type 1: A liquid stream RqiL from the tray j to tray q. (Liquid flowing downward)

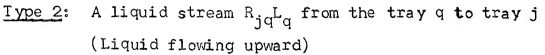
$$\overline{J} = \begin{bmatrix} B_1 & C_1 \\ A_2 & B_2 & C_2 \\ & & & & \\ & & & A_j & B_j & C_j \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ &$$

where

$$\overline{\overline{A}}_{q,j} = \begin{bmatrix} R_{qj} & \overline{\overline{I}}_c & \overline{\overline{O}}_c & \overline{\overline{O}}_c \\ \overline{\overline{O}}_c & \overline{\overline{O}}_c & \overline{\overline{O}}_c \\ \overline{\overline{X}}_c & \overline{\overline{O}}_c & X \end{bmatrix}$$
Structure: similar to the tridiagonal A blocks

Position: below the tridiagonal

Structure: similar to the tri-



where.

 $\overline{\overline{C}}_{j,q} = \begin{bmatrix} R_{j,q}\overline{\overline{L}}_c & \overline{\overline{O}}_c & \overline{\overline{O}}_c \\ \overline{\overline{O}}_c & \overline{\overline{\overline{O}}}_c & \overline{\overline{O}}_c \\ \overline{\overline{X}}_c & \overline{\overline{O}}_c & X \end{bmatrix}$ Structure: similar to the tridiagonal A-blocks

Position: above the tridiagonal

A vapor stream $R_{iq}V_{q}$ from tray q to tray j Type 3: (Vapor flowing upward)

$$\bar{J} = \begin{bmatrix} B_1 & C_1 \\ A_2 & B_2 & C_2 \\ & & & & \\ & & A_j & B_j & C_j & & & \\ & & & A_q & B_q^* & C_q \\ & & & & & \\ & & & & A_N & B_N \end{bmatrix}$$
 and

$$\overline{\overline{C}}_{j,q} = \begin{bmatrix} \overline{\overline{O}}_{c} & \mathbf{R}_{jq} & \overline{\overline{I}}_{c} & \overline{\overline{O}}_{c} \\ \overline{\overline{O}}_{c} & \overline{\overline{X}}_{c} & \overline{\overline{O}}_{c} \\ \overline{\overline{O}}_{c} & \overline{\overline{X}}_{c} & X \end{bmatrix}$$
Structure: similar to tridia-gonal C-blocks

Position: above the diagonal

A vapor stream R_{qi} V_i from tray j to tray q. Type 4:

and

$$C_{q,j} = \begin{bmatrix} \overline{0}_c & R_{qj} \overline{1}_c & \overline{0}_c \\ \overline{0}_c & \overline{X}_c & \overline{0}_c \end{bmatrix}$$
Structure: similar to tridiagonal c-blocks
Position: below the diagonal

The reciprocal streams in an arrangement of columns cause more than one off-tridiagonal blocks to occurs in the Jacobian, one offdiagonal element per single stream. reciprocal streams may be handled by breaking them up into

single streams and the position and structure all offtridiagonal blocks can be determined by identifying its type as discussed earlier.

Consider a reciprocal stream between two trays, a liquid stream $R_{jq}L_{q}$, flowing from tray q to tray j and, a vapor stream $R_{qj}V_j$, flowing from tray j to tray q. The liquid and vapor streams may be identified as type 2 and type 4 individually, and thus the structure of the Jacobian is:

where,
$$A_{q,j} = \begin{bmatrix} \overline{\overline{O}}_c & R_{qj} \overline{\overline{I}}_c & \overline{\overline{O}}_c \\ \overline{\overline{O}}_c & \overline{\overline{X}}_c & \overline{\overline{O}}_c \\ \overline{\overline{O}}_c & \overline{\overline{X}}_c & X \end{bmatrix}$$
Structure: similar to c-block.
Position: below the diagonal (corresponds to the vapor stream)

Structure: similar to c-blocks

an d

$$C_{j,q} = \begin{bmatrix} R_{j,q} \overline{\overline{I}}_c & \overline{\overline{O}}_c & \overline{\overline{O}}_c \\ \overline{\overline{O}}_c & \overline{\overline{O}}_c & \overline{\overline{O}}_c \\ \overline{\overline{V}}_c & \overline{\overline{O}}_c & X \end{bmatrix}$$
Structure: similar to A-blocks
Position: above the diagonal (Corresponds to the liquid stream)

(Corresponds to the liquid stream)

Thus, the position of an off-diagonal element depends only on the direction of the corresponding stream. In the Jacobian matrix, it is positioned in the row corresponding to the tray into which the stream enters. and the column corresponds to the tray from which it exits. Therefore, if the stream flows upward its position would be above the tridiagonal, and below the tridiagonal if the stream flows downwards, in a given arrangement of columns.

The structure of the off-diagonal elements may be determined only by knowing whether it is a liquid or a vapor stream. An offdiagonal element corresponding to a vapor stream has a structure similar to that of a tridiagonal C-block, and similar to that of a tridiagonal A-block if it is a liquid stream.

The detailed calculations of the elements of the tridiagonal and offdiagonal elements have been presented in the Appendix A.

Modelling of Condensers and Reboilers 2.3

Although the condensers and reboilers are treated like trays, the generalized tray model, discussed in

Section 2.1, cannot directly be applied since in these cases a few streams are missing and some additional specifications are required. In a condenser the liquid stream l_{j-1} , and in a reboiler the vapor stream v_{j+1} are not present. Moreover, there may be an additional specification for each condenser and reboiler (These may be specified in various ways, 13 of them are included in the Chapter 4).

We assume that there are no interlinking streams present in the condensers and reboilers, and the heat-duties are specified. (Refer figure 3 for various types of condensers and reboilers).

Partial Condensers:

Component Mass Balances:

$$M_{j,i} = r_{vj+1} v_{j+1,i} - (1+s_j)v_{j,i} - (1+s_j)l_{j,i}$$

$$1 \quad i \quad C \qquad (2.25)$$

Equilibrium Relationships:

$$Q_{j,i} = \frac{\eta_{j} K_{j,i}}{L_{j}} 1_{j,i} - \frac{v_{j,i}}{V_{j}} + \frac{(1 - \eta_{j})}{V_{j+1}} v_{j+1,i}$$

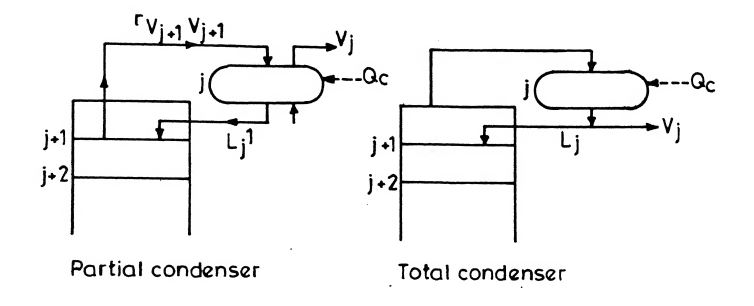
$$1 \le i \le C \qquad (2.26)$$

Enthalpy Balance:

$$E_{j} = r_{vj+1} \qquad \sum_{i=1}^{c} H_{j+1,i} \quad v_{j+1,i} - (1+S_{j}) \quad \sum_{i=1}^{c} H_{j,i} \quad v_{j,i} - (1+S_{j}) \quad \sum_{i=1}^{c} h_{j,i} \quad 1_{j,i} + Q_{c}$$

$$(2.27)$$

where, Qc is the condenser-heat-duty.



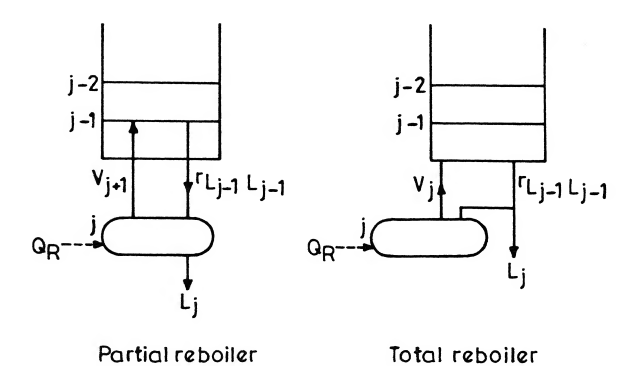


Fig.3

Total Condensers:

In the case of total condensers both the streams leaving the condenser are liquid streams, and have the same composition. We use the notation V_j for the distillate (which is liquid).

Component mass balances:

$$M_{j,i} = r_{vj+1} v_{j+1,i} - (1 + S_j)v_{j,i} - (1 + S_j)l_{j,i}, 1 \le i \le C$$
(2.28)

Equilibrium relationships: (Since there is no equilibrium, these are replaced by x_{j,i} = y_{j,i} equations)

$$Q_{j,i} = L_j V_{j,i} - V_j l_{j,i}, 1 \le i \le C$$
 (2.29)

Enthalpy Balance:

$$E_{j} = r_{vj+1} \sum_{i=1}^{c} H_{j+1,i} v_{j+1,i} - \sum_{i=1}^{c} h_{j,i} [(1+S_{j})v_{j,i} + (1+S_{j})l_{j,i}] + Q_{c}$$

$$(2.30)$$

Partial Reboilers:

Component mass balances:

$$M_{j,i} = r_{L_{j-1}} l_{j-1,i} - (1+s_j)l_{j,i} - (1+s_j)v_{j,i}, 1 \le i \le C$$
(2.31)

Equilibrium Relationships:

$$Q_{j,i} = \frac{K_{j,i}}{L_{j}} l_{j,i} - \frac{V_{j,i}}{V_{j}}, 1 \le i \le C$$
 (2.32)

Enthalpy Balance:

$$E_{j} = r_{Lj-1} \qquad \sum_{i=1}^{c} h_{j-1,i} l_{j-1} - (1+s_{j}) \sum_{i=1}^{c} h_{j,i} l_{j,i} - (1+s_{j}) \sum_{i=1}^{c} H_{j,i} v_{j,i} + Q_{R}$$
 (2.33)

where Q_R is the reboiler-heat-duty:

Total Reboiler:

In the case of a total reboiler a part of the L_{j-1} stream is drawn out as the bottom stream L_j , and the remaining is vaporized and fed to the column as V_j . Therefore, the bottom stream L_j is at the same temperature as that of L_{j-1} , and compositions of L_j and V_j streams are the same.

Component mass balances:

$$M_{j,i} = r_{Lj-1} l_{j-1,i} - (1+s_j)l_{j,i} - (1+s_j)v_{j,i} 1 \le i \le C$$
(2.34)

Equilibrium Relationships: (Since there is no equilibrium, these are replaced by x_{j,i} = y_{j,i} equations)

$$Q_{j,i} = L_j v_{j,i} - V_j l_{j,i} \quad 1 \le i \le C$$
 (2.35)

Enthalpy Balance:

$$E_{j} = \sum_{i=1}^{c} h_{j-1,i} [r_{Vj-1} l_{j-1,i} - (1+s_{j}) l_{j,i}] - (1+s_{j}) \sum_{i=1}^{c} H_{j,i} v_{j,i} + Q_{R}$$
(2.36)

There may be various other specifications besides the condenser and reboiler heat-duty specifications, in which case the enthalpy balances discussed above will no longer be applicable. These are therefore replaced by some other suitable equations which take into account the given specifications. A variety of specifications and alternative discrepancy functions to replace E_j are presemted in Chapter 4.

With the replacement of the discrepancy function E_j, its partial derivatives will also change. Therefore the last rows of the submatrices (B & C in case of condensers and A & B for reboilers) must be modified accordingly.

2.4 Method of Solution

In case of a single column problem with no bypassing or pump arounds, the Jacobian has a completely block-tridiagonal structure, and the well known Thomas-alogorithm may be applied to obtain the solution.

Thomas Algorithm:

Forward Substitution:

Step 1:
$$\overline{\overline{P}}_1 \leftarrow (\overline{\overline{B}}_1)^{-1} \overline{\overline{C}}_1$$

Step 2:
$$\overline{Q}_1 + (\overline{B}_1)^{-1} \overline{F}_1$$

For stages j, from 2 to (N-1)

Step 3:
$$\overline{P}_j + (\overline{B}_j - \overline{A}_j \overline{P}_{j-1})^{-1} \overline{C}_j$$

Step 4:
$$\bar{Q}_{j} * (\bar{B}_{j} - \bar{A}_{j} \bar{P}_{j-1})^{-1} (\bar{F}_{j} - \bar{A}_{j} \bar{Q}_{j-1})$$

For stage N

Step 5:
$$\bar{Q}_N \sim (\bar{B}_N - \bar{A}_N \bar{P}_{N-1})^{-1} (\bar{F}_N - \bar{A}_N \bar{Q}_{N-1})$$

Backward Substitution:

Step 6:
$$\Delta \vec{X}_N \leftarrow \vec{Q}_N$$

For stages j, from (N-1) to 1

Step 7:
$$\Delta \bar{X}_j + (\bar{Q}_j - \bar{\bar{P}}_j \Delta \bar{X}_{j+1})$$

If in a column bypass or pump-arounds are present, or if an arrangement of interlinked column is considered, the Jacobian does not have a strictly block-tridiagonal structure. Instead, a few off-tridiagonal blocks occur in the Jacobian, and consequently, the conventional Thomas algorithm cannot directly be applied.

A modification of the Thomas algorithm was developed by Hofeling and Seader [2], to solve a system in which the Jacobian has a few off-tridiagonal blocks:

Hofeling and Seader [2], have taken a specific problem to illustrate how the off-tridiagonal blocks could be handled by the modified Thomas algorithm.

Seader [6], justified that using the same principles of the modified Thomas algorithm, any arrangement of the off-tridiagonal blocks in a Jacobian could be handled.

There are a numerous ways, in which the off-tridiagonal blocks may appear in the Jacobian, varying from
problem to problem. Moreover, for a given system of interlinked columns, there may be a large number of different
possible arrangement of these off-tridiagonal blocks in
the Jacobian, depending upon the ordering of the columns or
column units (refer Hildalgo-Seader[4]). Although the ways
to handle these blocks, by the modified Thomas algorithm,
are the same in principle, it is difficult to develop a
step by step generalized algorithm to handle all possible
arrangements of the off-tridiagonal blocks in a Jacobian.

Consider the system of interlinked columns, in which there are two absorbers A_1 and A_2 of 10 plates each, and two distillation columns D_1 and D_2 having 15 and 12 plates respectively. These columns are interlinked as shown in the Figure 6. The Jacobian-matrix, for this system, has six

off-tridiagonal blocks, as shown in Figure 7.

We employ the modified Thomas algorithm to solve this system. Various steps of the forward substitution and finally of the backward substitution to obtain a correction vector are presented here.

Modified Thomas Algorithm:

Step 1: row 1,
$$P_1 + B_1^{-1} C_1$$
; $P_{1,20} + B_1^{-1} C_{1,20}$; $Q_1 + B_1^{-1} F_1$

Step 2: rows j, where
$$2 \le j \le 9$$
, $B_j = (B_j - A_j P_{j-1})^{-1}$;
$$P_j = B_j C_j; P_{j,20} = -B_j A_j P_{j-1,20}; Q_j = B_j (F_j - A_j Q_{j-1})$$

Step 3: row 10,
$$B_{10} - (B_{10} - A_{10}P_{9})^{-1}$$

$$P_{10} \leftarrow \overline{0}$$
; $P_{10,20} \leftarrow -B_{10}A_{10}P_{9,20}$; $Q_{10} \leftarrow B_{10}(F_{10}-A_{10}Q_{9})$

Step 4: row 11,
$$P_{11} + B_{11}^{-1} C_{11}$$
; $P_{11,47} + B_{11}^{-1} C_{11,47}$; $P_{11,20} + \overline{O}$; $Q_{11} + B_{11}^{-1} F_{11}$

Step 5: row j, where
$$12 \le j \le 18$$
, $B_{j} + (B_{j} - A_{j}P_{j-1})^{-1}$

$$P_{j} + B_{j}C_{j}; P_{j,20} + \overline{0}; P_{j,47} + -B_{j}A_{j}P_{j-1,47};$$

$$Q_{j} + B_{j}(F_{j} - A_{j}Q_{j-1})$$

itep 6: row 19,
$$B_{19} - (B_{19} - A_{19} P_{18})^{-1}$$

$$P_{19} + B_{19}C_{19}; \quad P_{19,47} + -B_{19}A_{19}P_{18,47};$$

$$Q_{19} + B_{19}(F_{19} - A_{19}Q_{18})$$
Step 7: row 20, set $\beta_1 + Q_{10}$; set $\beta_2 + P_{10,20}$;
iterate on $\beta_1 + Q_1$; from $j = 9$ to $j = 1$;
iterate on $\beta_2 + P_{j,20} - P_j\beta_2$, from $j = 9$ to $j = 1$;
$$B_{20} + (B_{20} - A_{20}P_{19} - A_{20,1}\beta_2)^{-1}$$

$$P_{20} + \overline{0}; \quad P_{20,47} + -B_{20}A_{20}P_{19,47};$$

$$Q_{20} + B_{20}(F_{20} - A_{20}Q_{19} - A_{20,1}\beta_1)$$
Step 8: row 21, $P_{21} + B_{21}^{-1} C_{21}; Q_{21} + B_{21}^{-1} F_{21}; P_{21,47} + \overline{0}$
Step 9: row 22, $B_{22} + (B_{22} - A_{22}P_{21})^{-1};$

$$P_{22} + B_{22}C_{22}; \quad P_{22,47} + \overline{0}; \quad Q_{22} + B_{22}(F_{22} - A_{22}Q_{21})$$
Step 10: row 23, $B_{23} + (B_{23} - A_{23}P_{22})^{-1}$

$$P_{23} + B_{23}C_{23}; \quad P_{23,47} + B_{23}A_{23,10}P_{10,20}P_{20,47}$$

$$Q_{23} + B_{23}[F_{23} - A_{23}Q_{22} - A_{23,10}(Q_{10} - P_{10,20}Q_{20})]$$
Step 11: rows j , where $24 \le j \le 34$; $B_j + (B_j - A_jP_{j-1})^{-1}$

$$P_j + B_jC_j; \quad P_j, 47 + B_jA_jP_{j-1,47}; \quad Q_j + B_j (F_j - A_jQ_{j-1})$$
Step 12: row 35, $B_{35} + (B_{35} - A_{35}P_{34})^{-1}$

 $P_{35} + \overline{0}$; $P_{35,47} + -B_{35}A_{35}P_{34,47}$; $Q_{35} + B_{35}(F_{35}-A_{35}Q_{34})$

Step 13: row 36,
$$P_{36} * B_{36}^{-1} C_{36}$$
; $P_{36,47} * \bar{0}$; $Q_{36} * B_{36}^{-1} F_{36}$
Step 14: rows j, where $37 \le j \le 39$; $B_j * (B_j - A_j P_{j-1})^{-1}$
 $P_j * B_j C_j$; $P_{j,47} * \bar{0}$; $Q_j * B_j (F_j - A_j Q_{j-1})$
Step 15: row 40, $B_{40} * (B_{40} - A_{40} P_{39})^{-1}$
reset $\beta_1 * Q_{35}$; iterate on $\beta_1 * Q_j - P_j \beta_1$, from $j = 34$ to $j = 21$ reset $\beta_2 * P_{35,47}$; iterate on $\beta_2 * P_{j,47} - P_j \beta_2$, from $j = 34$ to $j = 23$ $\beta_2 * P_{21} P_{22} \beta_2$
 $P_{40} * B_{40} C_{40}$; $P_{40,47} * - B_{40} A_{40,21} \beta_1$)
Step 16: row 41, $B_{41} * (B_{41} - A_{41} P_{40})^{-1}$; $P_{41} * B_{41} C_{41}$; $P_{41,47} * [-B_{41} (A_{41} P_{40,47} + A_{41,20} P_{20,47})]$
 $Q_{41} * B_{41} (F_{41} - A_{41} Q_{40} - A_{41,20} Q_{20})$
Step 17: rows j, where $42 \le j \le 45$, $B_j * (B_j - A_j P_{j-1})^{-1}$
 $P_j * B_j C_j$; $P_{j,47} * - B_j A_j P_{j-1} A_j C_j + B_j (F_j - A_j Q_{j-1})$
Step 18: row 46, $B_{46} * (C_{46} - A_{46} P_{45,47})$; $Q_{46} * B_{46} (F_{46} - A_{46} Q_{45})$
Step 19: row 47, $Q_{47} * (B_{47} - A_{47} P_{46})^{-1} (F_{47} - A_{47} Q_{46})$

Back Substitution:

Step 1:

Step 1: row 47,
$$X_{47} \leftarrow Q_{47}$$

Step 2: row 46, $X_{46} \leftarrow Q_{46} - P_{46}X_{47}$
Step 3: rows j, from j = 45 to j = 40
 $X_{j} \leftarrow Q_{j} - P_{j}X_{j+1} - P_{j,47}X_{47}$
Step 4: rows j, from j = 39 to j = 36
 $X_{j} \leftarrow Q_{j} - P_{j}X_{j+1}$
Step 5: row 35, $X_{35} \leftarrow Q_{35} - P_{35,47}X_{47}$
Step 6: rows j, from j = 34 to j = 23
 $X_{j} \leftarrow Q_{j} - P_{j}X_{j+1} - P_{j,47}X_{47}$
Step 7: rows j, from j = 22 and j = 21
 $X_{j} \leftarrow Q_{j} - P_{j}X_{j+1}$
Step 8: row 20, $X_{20} \leftarrow Q_{20} - P_{20,47}X_{47}$
Step 9: rows j, from j = 19 to j = 11
 $X_{j} \leftarrow Q_{j} - P_{j}X_{j+1} - P_{j,47}X_{47}$

Step 10: row 10, $X_{10} + Q_{10} - P_{10.20} X_{20}$

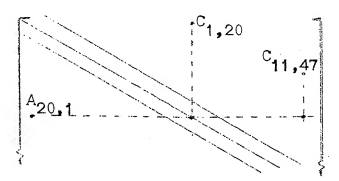
 $X_{i} + Q_{i} - P_{j} X_{i+1} - P_{j,20} X_{20}$

Step 11: rows j, from j = 9 to j = 1

It may be noted here that the calculations for eliminating the lower off-tridiagonal blocks is different for the blocks A20,1, A23,10 and A40,21 and A41,20. Each case is discussed

here separately:

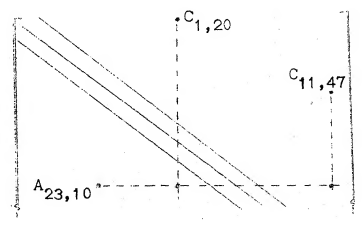
Row 20: Block $A_{20,1}$: The structure of Jacobian between rows 1 and 20 is as under:



Here, the row of $A_{20,1}$ and column of $C_{1,20}$ intersect on the diagonal, whereas those of $A_{20,1}$ and $C_{11,47}$ intersect at a point above the diagonal.

The calculation procedure for this type of situation is included in the step 7. Since the $\rm C_{20}$ block is zero, $\rm P_{20}$ block is also zero.

Row 23: Block A_{23,10}: The structure of the Jacobian up to row 23, is as follows:



In this case, the row of $A_{23,10}$ and the column of $C_{1,20}$ intersect at a point below the diagonal, whereas those of $A_{23,10}$ and $C_{11,40}$, above the diagonal.

The calculation procedure for this kind of situation is different from the previous case. Refer step 10. (It may be pointed out here that since C_{10} and C_{20} are zero therefore the blocks of rows between 10 and 20 and those of rows between 20 to 23 do not appear in the calculations). Please refer Figure 7.

If the blocks C_{10} and C_{20} were not zeros, then the step 10 would have been:

row 23, set $\beta_1 = Q_{19}$,

iterate on $\beta_1 + Q_j - P_j \beta_1$, from j = 18 to j = 10; set $\beta_2 + P_{19,47}$.

iterate on $\beta_2 + P_{j,47} - P_{j\beta_2}$, from j = 18 to j = 11; $\beta_2 + P_{10}\beta_2$;

set β 3 * P_{19,20}

iterate on $\beta_3 + P_{j,20} - P_{j}\beta_2$, from j = 18to j = 11; $\beta_3 + P_{10}\beta_3$;

set α₁ • Q₂₂

iterate on $\alpha_1 \leftarrow Q_j - P_j \alpha_1$, from j = 21 to j = 20set $\alpha_2 \leftarrow P_{22,47}$

iterate on $\alpha_2 = P_{j,47} - P_j \alpha_2$, from j = 21 to j = 20 set $\alpha_3 = P_{22}$

iterate on
$$\alpha_3 + P_{j}\alpha_3$$
, from $j = 21$ to $j = 20$

$$B_{23} + (B_{23} - A_{23}P_{22} - A_{23,10}\beta_3\alpha_3)^{-1}$$

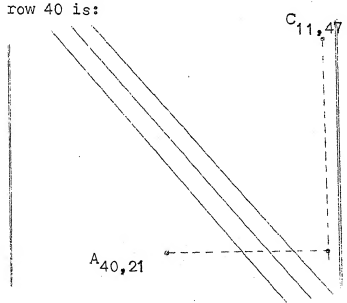
$$P_{23} + B_{23}C_{23};$$

$$P_{23,47} + B_{23}[A_{23,10}(\beta_2 - \beta_3\alpha_2) - A_{23}P_{22,47}]$$

$$Q_{23} + B_{23}[F_{23} - A_{23}Q_{22} - A_{23,10}(\beta_1 + \beta_3\gamma_1)]$$

Thus, the calculation procedure for this type of situation, becomes some what more complicated than the one discussed earlier.

Row 40: Block A40,21: The structure of the Jacobian upto



Here, the row of $A_{40,21}$ and the column of $C_{11,47}$ intersect at a point above the diagonal and there are no other off-tridiagonal blocks between the rows 21 and 39 after the forward substitution. Also the block C_{21} is non-zero.

The calculation procedure for this situation is described in the step 15.

Row 41: Block A41.20°

Since this situation is similar to that of the row 40, the calculation procedure followed is the same. However, it gets simplified to step 16 since the block C_{20} is zero, and therefore the blocks between rows 20 and 40 do not enter in the computations.

Thus, the numerous possible arrangements of off-tri-diagonal blocks which require different computational efforts, and the presence of some null blocks which may change the computations substantially, make it extremely difficult to develop a 'universal' algorithm which can handle all possible arrangements.

CHAPTER 3

SPARSITY EXPLOITATION

The conventional and modified Thomas algorithms exploit the sparsity of the Jacobian matrix which is highly sparse, and has a completely or almost-tridiagonal-band structure.

Moreover, the submatrices within the Jacobian are also sparse (Particularly A and C), and have a definite structure (refer 2.19, 2.20 and 2.21). The presence of zero and identity blocks within the submatrices and their definite structures, allow a further exploitation of the sparsity, which consequently results in a substantial reduction of both computational and storage requirements.

In this chapter, the exploitation of sparsity within the submatrices along with saving in storage and operationcount, is presented.

As is customary, only the operations involving multiplications and divisions are counted, neglecting those
involving addition and subtraction since the time consumed
in these is comparatively negligible.

To take the advantage of the sparsity and structure of the submatrices, computation and storage involving zero and unity blocks is avoided, which results in a substantial reduction in the CPU-time and memory-requirement on a computer.

Listed below are the various operations involving multiplication and inversion of submatrices, while exploiting their sparsity and structures, and comparison of the operation counts.

It may be pointed out that the P-blocks have two different structures viz. $P_{\rm V}$ and $P_{\rm I}$, which are as under:

$$\mathbf{P}^{V} = \begin{bmatrix} \overline{\mathbf{o}}_{\mathbf{c}} & \overline{\mathbf{x}}_{\mathbf{c}} & \overline{\mathbf{x}}_{\mathbf{c}} \\ \overline{\mathbf{o}}_{\mathbf{c}} & \overline{\mathbf{x}}_{\mathbf{c}} & \overline{\mathbf{x}}_{\mathbf{c}} \\ \overline{\mathbf{o}} & \overline{\mathbf{x}}_{\mathbf{c}} & \mathbf{x} \end{bmatrix} \quad ; \quad \mathbf{P}^{L} = \begin{bmatrix} \overline{\mathbf{x}}_{\mathbf{c}} & \overline{\mathbf{o}}_{\mathbf{c}} & \overline{\mathbf{x}}_{\mathbf{c}} \\ \overline{\mathbf{x}}_{\mathbf{c}} & \overline{\mathbf{o}}_{\mathbf{c}} & \overline{\mathbf{x}}_{\mathbf{c}} \\ \overline{\mathbf{x}}_{\mathbf{c}} & \overline{\mathbf{o}}_{\mathbf{c}} & \mathbf{x} \end{bmatrix}$$

Also, $B^{I} = B^{-1}$ or $(B-A P^{V})^{-1}$, where both have the same structure

$$= \begin{bmatrix} \overline{\mathbf{x}}_{\mathbf{c}} & \overline{\mathbf{x}}_{\mathbf{c}} & \overline{\mathbf{x}} \\ \overline{\mathbf{x}}_{\mathbf{c}} & \overline{\mathbf{x}}_{\mathbf{c}} & \overline{\mathbf{x}} \\ \overline{\mathbf{x}}_{\mathbf{c}} & \overline{\mathbf{x}}_{\mathbf{c}} & \mathbf{x} \end{bmatrix}$$

Operation 1: P^V ► B^IC

$$\begin{bmatrix} \overline{o} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \end{bmatrix} \leftarrow \begin{bmatrix} \overline{x}_{c} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{x}_{c} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{x}_{c} & \overline{x}_{c} & \overline{x}_{c} \end{bmatrix} \begin{bmatrix} \overline{o}_{c} & \mathbf{r} & \overline{\mathbf{I}}_{c} & \overline{o}_{c} \\ \overline{o}_{c} & \overline{x}_{c} & \overline{o}_{c} \\ \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \end{bmatrix}$$

$$P_{\mathbf{i},k+c}^{V} \leftarrow \mathbf{r} \quad B_{\mathbf{i},k}^{I} + \sum_{m=c+1}^{2c+1} B_{\mathbf{i},m} \quad C_{m,k+c} \quad 1 \leq i \leq 2c+1 \\ 1 \leq k \leq c$$

$$P_{i,2c+1}^{V}$$
 + $P_{i,2c+1}^{I}$ $P_{i,2c+1}^{C}$ + $P_{i,2c+1}^{C$

Operation 2: PV + BIA

$$\begin{bmatrix} \overline{X}_{c} & \overline{O}_{c} & \overline{X}_{c} \\ \overline{X}_{c} & \overline{O}_{c} & \overline{X}_{c} \\ \overline{X}_{c} & \overline{O}_{c} & \overline{X}_{c} \\ \overline{X}_{c} & \overline{O}_{c} & X \end{bmatrix} - \begin{bmatrix} \overline{X}_{c} & \overline{X}_{c} & \overline{X}_{c} \\ \overline{X}_{c} & \overline{X}_{c} & \overline{X}_{c} \\ \overline{X}_{c} & \overline{X}_{c} & X \end{bmatrix} \begin{bmatrix} r\overline{I}_{c} & \overline{O}_{c} & \overline{O}_{c} \\ \overline{O}_{c} & \overline{O}_{c} & \overline{O}_{c} \\ \overline{X}_{c} & \overline{O}_{c} & X \end{bmatrix}$$

$$P_{i,k+c}^{V} + r_{i,k}^{I} + B_{i,2c+1}^{I} A_{2c+1,k} + A_$$

^{*} Standard Operation Count

⁺ Improved Operation Count.

$$\begin{bmatrix} \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{o}_{c} & \overline{o}_{c} & \overline{o}_{c} \\ \overline{o}_{c} & \overline{o}_{c} & \overline{o}_{c} \end{bmatrix} + \begin{bmatrix} \overline{r}_{c} & \overline{o}_{c} & \overline{o}_{c} \\ \overline{o}_{c} & \overline{o}_{c} & \overline{o}_{c} \\ \overline{x}_{c} & \overline{o}_{c} & \overline{x} \end{bmatrix} \begin{bmatrix} \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \end{bmatrix}$$

$$R_{i,k+c} \leftarrow r^{p_{i,k+c}^{V}}$$

$$1 \le i \le c$$

$$1 \le k \le c+1$$

R_{2c+1,k+c}
$$\leftarrow$$
 A_{2c+1,2c+1} P_{2c+1,k+c} $+$ $\sum_{m=1}^{c}$ A_{2c+1,m}P_{m,k+c} $+$ $\sum_{m=1}^{c}$ A_{2c+1,m}P_{m,k+c} $+$ $+$ $\sum_{m=1}^{c}$ A_{2c+1,m}P_{m,k+c}

S.O.C. =
$$8C^3 + 12C^2 + 6C + 1$$

I.O.C. = $2C^2 + 2C + 1$
Saving = $8C^3 + 10C^2 + 4C$

Operation 4: $R \leftarrow A P^{L}$

$$\begin{bmatrix} \overline{X}_{c} & \overline{O}_{c} & \overline{X}_{c} \\ \overline{O}_{c} & \overline{O}_{c} & \overline{O}_{c} \\ \overline{X}_{c} & \overline{O}_{c} & X \end{bmatrix} \leftarrow \begin{bmatrix} \overline{I}_{c} & I\overline{O}_{c} & \overline{O}_{c} \\ \overline{O}_{c} & \overline{O}_{c} & \overline{O}_{c} \\ \overline{X}_{c} & \overline{O}_{c} & X \end{bmatrix} \begin{bmatrix} \overline{X}_{c} & \overline{O}_{c} & \overline{X}_{c} \\ \overline{X}_{c} & \overline{O}_{c} & \overline{X}_{c} \\ \overline{X}_{c} & \overline{O}_{c} & X \end{bmatrix}$$

$$R_{i,k} + r \quad P_{i,k}^{L}$$
 $1 \le i \le c$ $1 \le k \le c+1$

$$R_{2c+1,k} + A_{2c+1,2c+1} P_{2c+1,k}^{L} + \sum_{m=1}^{c} A_{2c+1,m} P_{m,k}^{L} \le k \le c$$
 $R_{2c+1,2c+1} + A_{2c+1,2c+1} P_{2c+1,2c+1}^{L} + \sum_{m=1}^{c} A_{2c+1,m} P_{m,2c+1}^{L}$
 $S.O.C. = 8C^3 + 12C^2 + 6C + 1$
 $I.O.C. = 2C^2 + 2C + 1$

Saving = $8C^3 + 10C^2 + 4C$

Operation 5:
$$R \leftarrow C P^V$$

$$\begin{bmatrix} \overline{o} & \overline{x}_c & \overline{x}_c \\ \overline{o}_c & \overline{x}_c & \overline{x}_c \\ \overline{o}_c & \overline{x}_c & \overline{x}_c \end{bmatrix} \leftarrow \begin{bmatrix} \overline{o}_c & r\overline{1}_c & \overline{o}_c \\ \overline{o}_c & \overline{x}_c & \overline{o}_c \\ \overline{o}_c & \overline{x}_c & \overline{x}_c \end{bmatrix} \leftarrow \begin{bmatrix} \overline{o}_c & r\overline{1}_c & \overline{o}_c \\ \overline{o}_c & \overline{x}_c & \overline{x}_c \\ \overline{o}_c & \overline{x}_c & \overline{x}_c \end{bmatrix}$$

$$R_{i,k+c} \leftarrow r P_{i+c,k+c}^V \qquad 1 \le i \le c, 1 \le k \le c+1$$

$$R_{i+c,k+c} \leftarrow \sum_{m=c+1}^{2c} C_{i+c,m} P_{m,k+c}^V \qquad 1 \le i \le c, 1 \le k \le c+1$$

$$R_{2c+1,k+c} \leftarrow \sum_{m=c+1}^{2c+1} C_{2c+1,m} P_{m,k+c}^V \qquad 1 \le k \le c+1$$

$$S.o.c. = 8c^3 + 12c^2 + 6C + 1$$

I.O.C. = $c^3 + 3c^2 + 3c + 1$

Saving = $7c^3 + 9c^2 + 3c$

$$\begin{bmatrix} \overline{X}_{\mathbf{c}} & \overline{\mathbf{o}}_{\mathbf{c}} & \overline{X}_{\mathbf{c}} \\ \overline{X}_{\mathbf{c}} & \overline{\mathbf{o}}_{\mathbf{c}} & \overline{X}_{\mathbf{c}} \end{bmatrix} + \begin{bmatrix} \overline{\mathbf{o}}_{\mathbf{c}} & \mathbf{r} \overline{\mathbf{I}}_{\mathbf{c}} & \overline{\mathbf{o}}_{\mathbf{c}} \\ \overline{\mathbf{o}}_{\mathbf{c}} & \overline{\mathbf{x}}_{\mathbf{c}} & \overline{\mathbf{o}}_{\mathbf{c}} \end{bmatrix} \begin{bmatrix} \overline{\mathbf{x}}_{\mathbf{c}} & \overline{\mathbf{o}}_{\mathbf{c}} & \overline{\mathbf{x}}_{\mathbf{c}} \\ \overline{\mathbf{x}}_{\mathbf{c}} & \overline{\mathbf{o}}_{\mathbf{c}} & \overline{\mathbf{x}}_{\mathbf{c}} \end{bmatrix} \begin{bmatrix} \overline{\mathbf{x}}_{\mathbf{c}} & \overline{\mathbf{o}}_{\mathbf{c}} & \overline{\mathbf{x}}_{\mathbf{c}} \\ \overline{\mathbf{x}}_{\mathbf{c}} & \overline{\mathbf{o}}_{\mathbf{c}} & \overline{\mathbf{x}}_{\mathbf{c}} \end{bmatrix}$$

$$R_{i,k} + r P_{i+c,k}^{L} \qquad 1 \le i \le c, 1 \le k \le c$$

$$R_{i,2c+1} + r P_{i+c,2c+1}^{L} \qquad 1 \le i \le c$$

$$R_{i+c,k} + \sum_{m=c+1}^{2c} C_{i+c,m} P_{m,k}^{L} \qquad 1 \le i \le c, 1 \le k \le c$$

$$R_{i+c,2c+1} + \sum_{m=c+1}^{2c} C_{i+c,m} P_{m,2c+1}^{L} \qquad 1 \le i \le c$$

$$R_{2c+1,k} + \sum_{m=c+1}^{2c+1} C_{2c+1,m} P_{m,k}^{L} \qquad 1 \le k \le c$$

$$R_{2c+1,k} + \sum_{m=c+1}^{2c+1} C_{2c+1,m} P_{m,k}^{L} \qquad 1 \le k \le c$$

$$R_{2c+1,2c+1} + \sum_{m=c+1}^{2c+1} C_{2c+1,m} P_{m,2c+1}^{L}$$

$$S.o.c. = 8c^{3} + 12c^{2} + 6c + 1$$

$$I.o.c. = c^{3} + 3c^{2} + 3c + 1$$

$$Saving = 7c^{3} + 9c^{2} + 3c$$

Operation 7:
$$P^{V} \leftarrow B^{I}R$$
, where $R = -AP^{V}$ (refer OP 3)

$$\begin{bmatrix} \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{o}_{c} & \overline{x}_{c} & x \end{bmatrix} \leftarrow \begin{bmatrix} \overline{x}_{c}^{\dagger} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{x}_{c} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{x}_{c} & \overline{x}_{c} & x \end{bmatrix} \begin{bmatrix} \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{o}_{c} & \overline{o}_{c} & \overline{o}_{c} \\ \overline{o}_{c} & \overline{x}_{c} & x \end{bmatrix}$$

$$P_{i,k+c}^{V} = B_{i,2c+1}^{I} R_{2c+1,k+c} + \sum_{m=1}^{c} B_{i,m}^{I} R_{m,k+c}$$

1 \(\text{i} \(\le 2c+1,1 \le k \le c+1 \)

S.O.C. =
$$8C^3 + 12C^2 + 6C + 1$$

I.O.C. = $2C^3 + 5C^2 + 4C + 1$
Saving = $6C^3 + 7C^2 + 2C$

Operation 8: $P^{L} \leftarrow B^{I}R$, where $R = -AP^{L}$ (refer OP 4)

$$\begin{bmatrix} \overline{x}_{c} & \overline{o}_{c} & \overline{x}_{c} \\ \overline{x}_{c} & \overline{o}_{c} & \overline{x}_{c} \\ \overline{x}_{c} & \overline{o}_{c} & x \end{bmatrix} \leftarrow \begin{bmatrix} \overline{x}_{c} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{x}_{c} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{x}_{c} & \overline{x}_{c} & x \end{bmatrix} \begin{bmatrix} \overline{x}_{c} & \overline{o}_{c} & \overline{x}_{c} \\ \overline{o}_{c} & \overline{o}_{c} & \overline{o}_{c} \\ \overline{x}_{c} & \overline{o}_{c} & x \end{bmatrix}$$

$$P_{i,k}^{L}$$
 $\leftarrow B_{i,2c+1}^{I}$ $R_{2c+1,k}$ $+$ $\sum_{m=1}^{c}$ $B_{i,m}$ $R_{m,k}$ $1 \le i \le 2c+1, 1 \le k \le c$

$$p_{i,2c+1}^{L}$$
 $p_{i,2c+1}^{R}$ p_{i

$$S.O.C. = 8C^3 + 12C^2 + 6C+1$$

$$I.O.C. = 2C^3 + 5C^2 + 4C+1$$

Saving =
$$6C^3 + 7C^2 + 2C$$

Operation 9:
$$P^{V} + B^{I}R$$
, where $R = C P^{V}$ (refer OP 5)

$$\begin{bmatrix} \overline{\overline{o}}_{c} & \overline{\overline{x}}_{c} & \overline{x}_{c} \\ \overline{\overline{o}}_{c} & \overline{\overline{x}}_{c} & \overline{x}_{c} \end{bmatrix} \leftarrow \begin{bmatrix} \overline{\overline{x}}_{c} & \overline{\overline{x}}_{c} & \overline{x}_{c} \\ \overline{\overline{x}}_{c} & \overline{\overline{x}}_{c} & \overline{x}_{c} \end{bmatrix} \begin{bmatrix} \overline{\overline{o}}_{c} & \overline{\overline{x}}_{c} & \overline{x}_{c} \\ \overline{\overline{o}}_{c} & \overline{\overline{x}}_{c} & \overline{x}_{c} \end{bmatrix} \begin{bmatrix} \overline{\overline{o}}_{c} & \overline{\overline{x}}_{c} & \overline{x}_{c} \\ \overline{\overline{o}}_{c} & \overline{\overline{x}}_{c} & \overline{x}_{c} \end{bmatrix}$$

$$P_{i,k+c}^{V}$$

$$\sum_{m=1}^{2c+1} B_{i,m}^{I} R_{m,k+c} \qquad 1 \leq i \leq 2c+1, 1 \leq k \leq c+1$$

$$5.0.C. = 8C^3 + 12C^2 + 6C + 1$$

$$I.O.C. = 4C^3 + 8C^2 + 5C + 1$$

Saving =
$$4C^3 + 4C^2 + C$$

Operation 10: $P^{L} \leftarrow B^{I}R$ where $R = C P^{L}$ (refer OP 6)

$$\begin{bmatrix} \overline{X}_{c} & \overline{o}_{c} & \overline{X}_{c} \\ \overline{X}_{c} & \overline{o}_{c} & \overline{X}_{c} \end{bmatrix} \leftarrow \begin{bmatrix} \overline{X}_{c} & \overline{X}_{c} & \overline{X}_{c} \\ \overline{X}_{c} & \overline{X}_{c} & \overline{X}_{c} \end{bmatrix} \begin{bmatrix} \overline{X}_{c} & \overline{o}_{c} & \overline{X}_{c} \\ \overline{X}_{c} & \overline{o}_{c} & \overline{X}_{c} \end{bmatrix} \begin{bmatrix} \overline{X}_{c} & \overline{o}_{c} & \overline{X}_{c} \\ \overline{X}_{c} & \overline{o}_{c} & \overline{X}_{c} \end{bmatrix}$$

$$P_{i,k}^{L} \leftarrow \sum_{m=1}^{2c+1} B_{i,m}^{I} R_{m,k}$$
 $1 \le i \le 2c+1, 1 \le k \le c$

$$P_{i,2c+1}^{L} + \sum_{m=1}^{2c+1} B_{i,m}^{I} R_{m,2c+1}$$
 1 $\leq i \leq 2c+1$

$$5.0.C. = 8C^3 + 12C^2 + 6C + 1$$

$$I.O.C. = 4C^3 + 8C^2 + 5C + 1$$

Saving =
$$4C^3 + 4C^2 + C$$

Operation 11 :
$$R \leftarrow p^{V} p^{L}$$

$$\begin{bmatrix} \overline{X}_{c} & \overline{\overline{o}}_{c} & \overline{X}_{c} \\ \overline{X}_{c} & \overline{\overline{o}}_{c} & \overline{X}_{c} \\ \overline{X}_{c} & \overline{\overline{o}}_{c} & X \end{bmatrix} + \begin{bmatrix} \overline{\overline{o}}_{c} & \overline{\overline{X}}_{c} & \overline{X}_{c} \\ \overline{\overline{o}}_{c} & \overline{\overline{X}}_{c} & \overline{X}_{c} \\ \overline{\overline{o}}_{c} & \overline{X}_{c} & X \end{bmatrix} \begin{bmatrix} \overline{X}_{c} & \overline{\overline{o}}_{c} & \overline{X}_{c} \\ \overline{\overline{X}}_{c} & \overline{\overline{o}}_{c} & \overline{X}_{c} \\ \overline{\overline{X}}_{c} & \overline{\overline{o}}_{c} & X \end{bmatrix}$$

$$R_{i,2c+1} \leftarrow \sum_{m=c+1}^{2c+1} P_{i,m}^{V} P_{m,2c+1}^{L} \qquad 1 \le i \le 2c+1$$

$$5.0.C. = 8C^3 + 12C^2 + 6C + 1$$

I.O.C. =
$$2C^3 + 5C^2 + 4C + 1$$

Saving =
$$6C^3 + 7C^2 + 2C$$

Operation 12: $R - P^{L} P^{L}$

$$\begin{bmatrix} \bar{X}_{c} & \bar{O}_{c} & \bar{X}_{c} \\ \bar{X}_{c} & \bar{O}_{c} & \bar{X}_{c} \\ \bar{X}_{c} & \bar{O}_{c} & \bar{X}_{c} \end{bmatrix} \leftarrow \begin{bmatrix} \bar{X}_{c} & \bar{O}_{c} & \bar{X}_{c} \\ \bar{X}_{c} & \bar{O}_{c} & \bar{X}_{c} \\ \bar{X}_{c} & \bar{O}_{c} & \bar{X} \end{bmatrix} \begin{bmatrix} \bar{X}_{c} & \bar{O}_{c} & \bar{X}_{c} \\ \bar{X}_{c} & \bar{O}_{c} & \bar{X}_{c} \\ \bar{X}_{c} & \bar{O}_{c} & \bar{X} \end{bmatrix} \begin{bmatrix} \bar{X}_{c} & \bar{O}_{c} & \bar{X}_{c} \\ \bar{X}_{c} & \bar{O}_{c} & \bar{X}_{c} \\ \bar{X}_{c} & \bar{O}_{c} & \bar{X}_{c} \end{bmatrix}$$

$$R_{i,k} \leftarrow P_{i,2c+1}^{L} P_{2c+1,k}^{L} + \sum_{m=1}^{c} P_{i,m}^{L} P_{m,k}^{L} 1 \le i \le 2c+1, 1 \le k \le c$$

$$R_{i,2c+1} \leftarrow P_{i,2c+1}^{L} P_{2c+1,2c+1}^{L} + \sum_{m=1}^{c} P_{i,m}^{L} P_{m,2c+1}^{L} 1 \le i \le 2c+1$$

$$S.O.C. = 8C^{3} + 12C^{2} + 6C + 1$$

$$I.O.C. = 2C^{3} + 5C^{2} + 4C + 1$$

Saving =
$$6C^3 + 7C^2 + 2C$$

Operation 13:
$$R \leftarrow P^{L_pV}$$

$$\begin{bmatrix} \overline{\overline{o}}_{c} & \overline{\overline{x}}_{c} & \overline{\overline{x}}_{c} \\ \overline{\overline{o}}_{c} & \overline{\overline{x}}_{c} & \overline{\overline{x}}_{c} \\ \overline{\overline{o}}_{c} & \overline{\overline{x}}_{c} & \overline{x} \end{bmatrix} + \begin{bmatrix} \overline{\overline{x}}_{c} & \overline{\overline{o}}_{c} & \overline{x}_{c} \\ \overline{\overline{x}}_{c} & \overline{\overline{o}}_{c} & \overline{x}_{c} \\ \overline{\overline{x}}_{c} & \overline{\overline{o}}_{c} & \overline{x} \end{bmatrix} \begin{bmatrix} \overline{\overline{o}}_{c} & \overline{\overline{x}}_{c} & \overline{x}_{c} \\ \overline{\overline{o}}_{c} & \overline{\overline{x}}_{c} & \overline{x}_{c} \\ \overline{\overline{o}}_{c} & \overline{\overline{x}}_{c} & \overline{x}_{c} \end{bmatrix}$$

 $1 \le i \le 2c+1$, $1 \le k \le c+1$

S.O.C. =
$$8C^3 + 12C^2 + 6C + 1$$

I.O.C. = $2C^3 + 5C^2 + 4C + 1$
Saving = $6C^3 + 7C^2 + 2C$

Operation 14: $R \leftarrow P^{V} P^{V}$

$$\begin{bmatrix} \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \end{bmatrix} \leftarrow \begin{bmatrix} \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \end{bmatrix} \begin{bmatrix} \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \end{bmatrix} \begin{bmatrix} \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \\ \overline{o}_{c} & \overline{x}_{c} & \overline{x}_{c} \end{bmatrix}$$

R_{i,k+c}
$$\stackrel{2c+1}{\underset{m}{\longrightarrow}} P_{i,m}^{V} P_{m,k+c}^{V}$$
 1 $\leq i \leq 2c+1$, 1 $\leq k \leq c+1$
S.O.C. = 8C³ + 12C² + 6C+1
I.O.C. = 2C³ + 5C² + 4C + 1
Saving = 6C³ + 7C² + 2C

$$\begin{bmatrix} \overline{X}_{c} \\ \overline{O}_{c} \\ X_{c} \end{bmatrix} \leftarrow \begin{bmatrix} \overline{r}_{c}^{T} & \overline{O}_{c} & \overline{O}_{c} \\ \overline{O}_{c} & \overline{O}_{c} & \overline{O}_{c} \\ \overline{X}_{c} & \overline{O}_{c} & X \end{bmatrix} \begin{bmatrix} \overline{X}_{c} \\ \overline{X}_{c} \\ \overline{X}_{c} \end{bmatrix}$$

$$s_i + rQ_i$$
, $1 \le i \le c$

$$s_{2c+1} - A_{2c+1,2c+1} - Q_{2c+1} + \sum_{m=1}^{c} A_{2c+1,m} - Q_{m}$$

$$S.O.C. = 4C^2 + 4C + 1$$

$$I.O.C. = 2C + 1$$

Saving =
$$4C^2 + 2C$$

Operation 16: S + CQ

$$\begin{bmatrix} \overline{X}_{c} \\ \overline{X}_{c} \\ X \end{bmatrix} + \begin{bmatrix} \overline{\overline{O}}_{c} & r\overline{\overline{I}}_{c} & \overline{\overline{O}}_{c} \\ \overline{\overline{O}}_{c} & \overline{X}_{c} & \overline{\overline{O}}_{c} \\ \overline{\overline{O}}_{c} & \overline{X}_{c} & X \end{bmatrix} \begin{bmatrix} \overline{X}_{c} \\ \overline{X}_{c} \\ \overline{X}_{c} \\ \overline{X}_{c} \end{bmatrix}$$

$$S_{i} + r Q_{i+c} \qquad 1 \le i \le c$$

$$S_{i} + r Q_{i+c} \qquad 0 \le i \le c$$

$$\mathbf{S}_{i+c}$$
 + $\sum_{m=c+1}^{2c}$ $C_{i+c,m}$ Q_m $1 \le i \le c$

$$S_{2c+1} + \sum_{m=c+1}^{2c+1} C_{2c+1,m} Q_{m}$$

$$5.0.C. = 4C^2 + 4C + 1$$

$$I.0.C. = C^2 + 2C + 1$$

Saving =
$$3C^2 + 2C$$

$$\begin{bmatrix} \overline{X}_{c} \\ \overline{X}_{c} \\ X \end{bmatrix} \leftarrow \begin{bmatrix} \overline{\overline{O}}_{c} & \overline{\overline{X}}_{c} \\ \overline{\overline{O}}_{c} & \overline{\overline{X}}_{c} \\ \overline{\overline{O}}_{c} & \overline{\overline{X}}_{c} \end{bmatrix} \begin{array}{c} \overline{X}_{c} \\ \overline{X}_{c} \\ \overline{X}_{c} \\ X \end{bmatrix}$$

$$S_{i}$$
 $\sum_{m=c+1}^{2c+1} P_{i,m}^{V} Q_{m}$ $1 \le i \le 2c+1$

$$S.O.C. = 4C^2 + 4C + 1$$

$$I.O.C. = 2C^2 + 3C + 1$$

Saving =
$$2C^2 + C$$

Operation 18: S + PLQ

$$\begin{bmatrix} \vec{x}_c \\ \vec{x}_c \\ x \end{bmatrix} \leftarrow \begin{bmatrix} \overline{\vec{x}}_c & \overline{\vec{o}}_c & \vec{x}_c \\ \overline{\vec{x}}_c & \overline{\vec{o}}_c & \vec{x}_c \\ \overline{\vec{x}}_c & \overline{\vec{o}}_c & x \end{bmatrix} \begin{bmatrix} \vec{x}_c \\ \overline{\vec{x}}_c \\ x \end{bmatrix}$$

$$S_{i} - P_{i,2c+1}^{L} Q_{2c+1} + \sum_{m=1}^{c} P_{i,m} Q_{m} \quad 1 \le i \le 2c+1$$

$$S_{0}C_{0} = 4C^{2} + 4c + 1$$

$$I.O.C. = 2C^2 + 3C + 1$$

Saving =
$$2C^2 + C$$

$$\begin{bmatrix} \overline{X}_{c} \\ \overline{X}_{c} \\ X \end{bmatrix} \leftarrow \begin{bmatrix} \overline{\overline{X}}_{c} & \overline{\overline{X}}_{c} & \overline{X}_{c} \\ \overline{\overline{X}}_{c} & \overline{\overline{X}}_{c} & \overline{X}_{c} \\ \overline{\overline{X}}_{c} & \overline{\overline{X}}_{c} & X \end{bmatrix} \begin{bmatrix} \overline{X}_{c} \\ \overline{X}_{c} \\ \overline{X}_{c} \end{bmatrix}$$

$$S_i \leftarrow \sum_{m=1}^{2c+1} B_{i,m}^{I} F_m \qquad 1 \leq i \leq 2c+1$$

5.0.C. =
$$4C^3 + 4C^2 + 1$$

I.0.C. = $4C^3 + 4C^2 + 1$
Saving = 0

Operation 20: Inversion of B or (B - AP)

The B and (B- AP) matrices have the same structure, which is as under:

$$\begin{bmatrix} -(1+s)\overline{1}_{c} & \overline{X}_{c} & \overline{X}_{c} \\ \overline{X}_{c} & \overline{X}_{c} & \overline{X}_{c} \\ \overline{X}_{c} & \overline{X}_{c} & X \end{bmatrix} = \begin{bmatrix} \overline{b}_{11} & \overline{b}_{12} \\ \overline{b}_{21} & \overline{b}_{22} \\ \overline{b}_{21} & \overline{b}_{22} \end{bmatrix}$$

Inversion by partitioning does not result in saving of the computational effort. It may be noted here that this matrix contains a cxc identity matrix, $I_{\rm c}$, and hence some saving in the operations may be achieved as demonstrated below.

We partition the matrix as shown above (one partion only)

Hence,
$$B^{-1}$$
 or $(B - AP)^{-1} = B^{I} = \begin{bmatrix} \overline{D}_{11} & \overline{D}_{12} \\ \overline{D}_{21} & \overline{D}_{22} \end{bmatrix}$
where,

order of
$$\overline{D}_{11} = cxc$$

order of $\overline{D}_{12} = cx(c+1)$
order of $\overline{D}_{21} = (c+1) \times c$
order of $\overline{D}_{22} = (c+1) \times (c+1)$

and,

$$\overline{\overline{D}}_{22} = (\overline{\overline{D}}_{22} - \overline{\overline{D}}_{21} (\overline{\overline{D}}_{11})^{-1} \overline{\overline{D}}_{12})^{-1}$$
(1)

$$\overline{\overline{D}}_{11} = (\overline{\overline{D}}_{11})^{-1} + (\overline{\overline{D}}_{11})^{-1} \overline{\overline{D}}_{12} \overline{\overline{D}}_{22} \overline{\overline{D}}_{21} (\overline{\overline{D}}_{21})^{-1}$$
 (2)

$$\bar{D}_{12} = -(\bar{D}_{11})^{-1} \bar{D}_{12} \bar{D}_{22}$$
 (3)

$$\bar{D}_{21} = -\bar{D}_{22} \bar{D}_{21} (\bar{D}_{11})^{-1}$$
 (4)

Since $(b_{11})^{-1} = -\frac{1}{(1+s)} \frac{1}{c}$, (only one division is needed to compute it), we need to perform only one inversion of the (c+1) x (c+1) in equation (1), to compute the B^{I} .

S.O.C. =
$$8C^3 + 12C^2 + 6C + 1$$

I.O.C. = $5C^3 + 12C^2 + 8C + 2$
Saving = $3C^3 - (2c+1)$
 $\approx 3C^3$

In the case of a single column problem with no pumparound or bypass streams, only the operations 1,3,15,17,19 and 20 are performed in the conventional Thomas algorithm. Moreover, the fractions $r_{\mathbf{V}j}$ and $r_{\mathbf{L}j}$ are unity for all the trays, since there are no interlinked-streams present. Therefore, in the operations 1,3 and 15 there is no need to multiply these fractions and a further reduction in the operation count is obtained. The improved operation count for these operations is presented below:

Improved operation count

Operation 1:
$$2C^3 + 3C^2 + 3C + 1$$

Operation 3: $C^2 + 2C + 1$

Operation 15: C+1

Saving in Storage Requirement:

In the conventional or modified Thomas algorithm, computations are performed stage by stage. Since the computations for any row j, require only tridiagonal and off-diagonal P-blocks of the previous j-1 rows, the A, B and C submatrices need not be stored, nor the identity blocks on the diagonal after the forward substitution. Thus, we need to store only the P-blocks. Since in both types of P-blocks, there are (2c+1) x c zero elements which need not be stored.

$$P^{V} = \begin{bmatrix} O_{c} & \overline{X}_{c} & \overline{X}_{c} \\ \overline{O}_{c} & \overline{X}_{c} & \overline{X}_{c} \\ \overline{O}_{c} & \overline{X}_{c} & X \end{bmatrix} \qquad P^{L} = \begin{bmatrix} \overline{X}_{c} & \overline{O}_{c} & \overline{X}_{c} \\ \overline{X}_{c} & \overline{O}_{c} & \overline{X}_{c} \\ \overline{X}_{c} & \overline{O}_{c} & X \end{bmatrix}$$

Thus, out of $(2c+1) \times (2c+1)$ elements of any P-block only $(2c+1) \times (c+1)$ are stored, and thereby almost a 50% further saving in storage results (Refer Chapter 6).

CHAPTER 4

SPECIFICATIONS

A multicomponent, multistage separation process problem must have the following specifications:

- (i) Number of trays
- (ii) Number of components
- (iii) Column pressure
- (iv) Complete specification of the feed (the feed rate, composition, thermal conditions, and the location of the feed tray).

In case of distillation, the location and type of every condenser and reboiler must also be specified.

In addition to these, some more specifications are needed to completely define the problem. These specifications may be made for the condensers, reboilers or for any other intermediate stages. It is important that the total number of these specifications must be equal to the total number of condensers and reboilers present in the system, otherwise the problem would become under or overspecified.

Specifications for Condensers and Reboilers:

In the Naphtali-Sandholm model, the specifications for condensers and reboilers are their heat duties, which are incorporated in the enthalpy balance discrepancy function, E;

For specifications other than the heat duties, the enthalpy balance in equations (2.27), (2.30), (2.33) or (2.36) shall no longer be applicable, and should therefore be replaced by the suitable discrepancy functions which takes into account the given specifications.

Some of these specifications and their corresponding discrepancy functions are presented in this chapter.

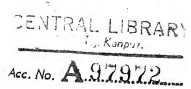
Specifications for the Intermediate Trays

When specifications are made at the intermediate stages other than condensers and reboilers, the heat-duties are no longer variable, and get fixed accordingly.

Consequently, the enthalpy balance discrepancy functions,

E; for the condensers and reboilers are no longer applicable.

Hofeling and Seader [2], have proposed that the inapplicable discrepancy function E_j , for a condenser or reboiler tray, may be replaced by a suitable discrepancy function, which incorporates the specification at the given intermediate stage. This replacement however generates an off-tridiagonal block in the Jacobian since the function E_j for a condenser or reboiler contains the variable(s) of an intermediate stage. The modified Thomas algorithm may be employed to handle the off-tridiagonal blocks.



Generation of the off-tridiagonal blocks results in additional computational and storage requirements, and should therefore be avoided, if possible. It is therefore proposed here, that the discrepancy function E_j for condenser or reboiler may be modified and made applicable (as in Spc. 1 and Spc. 2 below), and the E_j for the intermediate stage should be replaced by a suitable discrepancy function to incorporate the given specification at the same tray.

Listed below are some specifications and corresponding discrepancy functions which may be used to replace the inapplicable ones:

Spc. 1: If a specification is made at an intermediate stage instead of condenser, the condenser heat duty Q_c in equations 2.27 and 2.30 cannot be specified and should therefore be computed. Calculations of Q_c , and the modified E_j are as under: Partial Condenser: In a partial condenser, (which is represented by tray j), out of the r_{vj+1} , V_{j+1} moles of the vapor stream, $(1 + s_j)L_j$ moles are condensed (Figure 3). Therefore, Q_c may be computed by enthalpy considerations:

$$Q_c = -(1 + s_j) \sum_{i=1}^{c} (H_{j+1,i} - h_{j,i}) 1_{j,i}$$

and

$$E_{j} = \sum_{i=1}^{C} H_{j+1,i} (r_{Vj+1} v_{j+1,i} - (1+s_{j})l_{j,i}) - (1+s_{j}) \sum_{i=1}^{C} H_{j,i} v_{j,i}$$

Total condenser: In a total condenser, tray j, the whole r_{vj+1} v_{j+1} stream is condensed.

$$Q_{c} = -r_{v_{j+1}} \sum_{i=1}^{c} (H_{j+1,i} - h_{j,i})v_{j+1,i}$$

$$E_{j} = \sum_{i=1}^{c} h_{j,i} (r_{v_{j+1}} v_{j+1,i} - (1+s_{j})l_{j,i} - (1+s_{j})v_{j,i})$$

Spc. 2: If a specification is made at an intermediate stage instead of a reboiler, the reboiler heat duty Q_R in equation (2.33) and (2.36), cannot be specified and should therefore be computed.

Partial Reboiler: In a partial reboiler (which is represented by tray j), out of the r_{vj+1} L_{j-1} moles of the liquid stream, $(1 + S_j)V_j$ moles are vaporized (Figure 3). Q_R may be computed by enthalpy considerations:

$$Q_{R} = (1 + S_{j}) \sum_{i=1}^{C} (H_{j,i} - h_{j-1,i}) v_{j,i}$$

$$E_{j} = \sum_{i=1}^{C} h_{j-1,i} (r_{vj-1} l_{j-1,i} - (1 + S_{j})v_{j,i}) - \\ - (1 + S_{j}) \sum_{j=1}^{C} h_{j,i} l_{j,i}$$

Total Reboiler: In this case, out of r_{vj-1} L_{j-1} moles, $(1 + s_j)l_j$ moles are withdrawn and remaining $(1 + s_j)V_j$ are fed to the reboiler and are completely vaporized (Figure 3).

$$Q_{R} = (1 + S_{j}) \sum_{i=1}^{C} (H_{j,i} - h_{j-1,i})v_{j,i}$$

$$E_{j} = \sum_{i=1}^{C} h_{j-1,i} (r_{vj-1} l_{j-1,i} - (1+S_{j})l_{j,i}-(1+S_{j})v_{j,i})$$

- Spc. 3: Specification of the condenser-heat-duty, Q_c . As discussed earlier, (Please refer equations (2.27) and (2.30) for E_j).
- Spc. 4: Specification of the reboiler-heat-duty, $Q_{\mathbf{R}^{\bullet}}$ As discussed earlier (Please refer equations (2.33) and (2.36) for $E_{\mathbf{j}}$)
- Spc. 5: Specification of the Reflux Ratio, R = L/D for a condenser. (or R = (L/V) may be specified for an intermediate stage)

$$E_{j} = L_{j} - R V_{j}$$

Spc. 6: Specification of the Reboil Ratio, r = V/B for a reboiler. (or r = (V/L) may be specified for an intermediate stage) $E_j = V_j - r L_j$.

- Spc. 7: Specification of Temperature at any stage j $E_{j} = T_{j} T_{spec}.$

- Spc.10: Specification of molar vapor flow rate of component i at a stage $^{\circ}j$. $((v_{j,i})_{spec} = d_i = vapor key component flow rate, for a condenser tray). <math display="block">E_j = v_{j,i} (v_{j,i})_{spec}.$
- Spc.11: Specification of molar liquid flow rate of component i at a stage j. $((l_{j,i})_{spec} = b_i = liquid key component flow rate, for a reboiler tray). <math display="block">E_j = l_{j,i} (l_{j,i})_{spec}.$
- Spc.12: Specification of vapor mole fraction of component i at stage j. $((y_{j,i})_{spec.} = x_{d_i} = vapor key$ component composition for a condenser tray).
 - $E_j = v_{j,i} V_j(y_{j,i})_{spec}$
- Spc.13: Specification of liquid mole fraction of component i at stage j. $((x_{j,i})_{spec} = x_{b_i} = liquid key component composition for a reboiler tray)

 <math display="block">E_j = l_{j,i} L_j (x_{j,i})_{spec}$

Consider a single distillation column with 15 trays (including the condenser as tray 1 and reboiler as tray 15), and a 6-component mixture. Instead of condenser and reboiler heat-duties, the temperature of tray 9 and the flow rate of vapor leaving the tray 10 are specified.

As proposed by Hofeling and Seader in [2], the condenser enthalpy balance discrepancy function E_1 , is replaced by $E_1 = T_9 - (T_9)_{\text{specified}}$, and E_{15} of reboiler by $E_{15} = V_{10} - (V_{10})_{\text{specified}}$. The resulting Jacobian structure is shown in Figure 8, which may be solved using the modified Thomas algorithm.

Alternatively,we can modify the discrepancy functions E_1 and E_{15} as discussed in Spc. 1 and Spc. 2, and replace those of rows 9 and 10 by $E_9 = T_9 - (T_9)_{\text{specified}}$ and $E_{10} = V_{10} - (V_{10})_{\text{specified}}$, respectively. The resulting Jacobian matrix, in this case, has a completely tridiagonal structure with no off-tridiagonal elements. The conventional Thomas algorithm is employed here, and a substantial reduction in the computational efforts (3.877 times), and in memory requirements (2.786 times) is thereby realized.

CHAPTER 5

IMPLEMENTATION OF THE ALGORITHM

A computer program for solving the multicomponent separation process problems has been written in FORTRAN 10 and implemented on DEC 1090 system based on the algorithm described earlier. The program is kept in a file SEP.FOR.

The file SEP.FOR consists of a main program INTLNK and various subroutines and the tasks performed by each of them are listed below:

Main Program:

INTLNK:

The program may be considered to have two parts with different tasks. The first part reads the data, and computes the discrepancy function - vector. This part has been written for a general system of interlinked-columns and can handle any arbitrary arrangement of columns.

The second part includes the algorithm for solving a specific problem. As has been pointed out in Chapter 2, it is difficult to develop a generalized Hofeling-Seader algorithm which can handle all possible arrangements of the off-tridiagonal blocks in a Jacobian. Therefore this part must be rewritten for a given arrangement of the off-tridiagonal blocks.

Subroutines

ENL : Computes the liquid-molar-enthalpy for all components at a given stage (using a polynomial).

ENV : Computes the vapor-molar-enthalpy for all components at a given stage (using a polynomial).

FINDK: Computes the equilibrium K_{ij} values for all components for a given stage (using Antoine's constants (ideal-case)).

DKBYDX : Computes the derivatives $\frac{\partial k}{\partial x}$

DKBYDY : Computes the derivatives $\frac{\partial k}{\partial y}$

DKBYDT : Computes the derivatives $\frac{\partial k}{\partial T}$

DHVDT : Computes the derivatives $\frac{\partial H}{\partial T}$

DHLDT : Computes the derivatives $\frac{\partial h}{\partial T}$

CONBC : Computes the elements of the B & C submatrices for a partial or a total condenser.

REBAB : Computes the elements of the A & B submatrices for a partial or a total reboiler.

TRIA : Computes the elements of the A submatrix for a given tray.

TRIB : Computes the elements of the B submatrix for a given tray.

TRIC : Computes the elements of the C submatrix for a given tray.

OFFA: Computes the elements of the off-tridiagonal block having the same structure as that of an A-submatrix. This block is generated because of the liquid interlinked-stream.

OFFC: Computes the elements of the off-tridiagonal blocks having the same structure as that of C-matrix. This block is generated because of the vapor-interlinked stream.

PROB: Replaces the last row of the A, B and C submatrices to incorporate the given specification
for a stage j, where j may represent a condenser,
reboiler or an intermediate stage.

The following subroutines perform the various operations, which have been presented in the Chapter).

BCMUL : Operation 1

BAMUL : Operation 2

APVMUL : Operation 3

APLMUL : Operation 4

CPVMUL : Operation 5

CPLMUL : Operation 6

MBAPV : Operation 7

MBAPL : Operation 8

BCPV : Operation 9

BCPL	o •	Operation 10
PVPL	•	Operation 11
PLPL	0	Operation 12
PLPV	0	Operation 13
PVPV	9	Operation 14
AQMUL	•	Operation 15
CQMUL	•	Operation 16
QMPVX	0	Operation 17
QMPLX	9	Operation 18
BFMUL	° °	Operation 19
INVPRT	•	Operation 20 (This subroutine computes
		the inverse of a B or (B-AP) matrix
		of order (2c+1) x (2c+1))
MATINV	•	Computes the inverse of $(c+1)x(c+1)$
		matrix using Gauss Jordan method

(called by INVPRT).

with the maximum pivot strategy

After the input-data has been read, a check is conducted to find whether the problem is under or over specified, and if found so the execution is immediately terminated with an error message. Several checks are made at the various stages of execution, and if any inconsistency is detected, a suitable error message is printed out and the execution is stopped.

For a single column problem, the variable NOFF, which represents the number of off-tridiagonal blocks in the Jacobian, should be set to zero, and the program uses the conventional Thomas algorithm to solve the problem.

The thirteen different specifications for the condensers, reboilers and intermediate trays, which are presented in the Chapter 4, have been implemented in the program. For a given specification, a suitable code (1 through 13) is assigned to the variable LABEL and the specifications are handled following the approach discussed earlier. The subroutine PROB modifies the corresponding derivatives in the last row of the submatrices A, B, and C to incorporate the given specification.

In a system of interlinked columns a few vapor or liquid streams may be missing. This situation may arise in the cases where splitting and rearrangement of columns is done to obtain the "best-ordering" [4]. The locations of the trays for which a leaving-vapor stream is missing are stored in the JNO5V array, and those for the missing liquid streams in the JNO5L array. The missing streams are omitted in the calculations.

The fractions r_V and r_L (refer Chapter 2), are unity for the trays from which no interlinked stream is leaving,

and for all single column problems. Therefore these need not be included in the various multiplication operations such as operation 1,2 and 3 etc. The subroutines take care of this fact and a further saving in the operation count may be realized.

CHAPTER 6

RESULTS AND DISCUSSIONS

The exploitation of sparsity of the submatrices, while performing the matrix multiplications and inversions in the algorithms presented in Chapter 2, has resulted in a significant reduction in computational and storage requirements. The reduction in both, the operation count and the storage, for a variety of problems is presented in this chapter.

Problem 1: Consider a single absorption column having 8 stages and using a mixture of 14-components (For the complete specifications please refer [9]).

The problem was solved by applying the conventional Thomas algorithm, and a comparison of the operation counts is made in the Table 1.

A substantial reduction in the computation by a ratio of 3.02 was obtained by the sparsity exploitation, as only 183774 operations were performed instead of 555060.

Since only 3045 (= 7x29x14) elements need to be stored instead of 5887 (= 7x29x29), a saving of about 48% in the storage was realized.

This problem when implemented on DEC-10 took 4 iterations to converge to a tolerance limit of $9x10^{-4}$

(sum of squares of all the discrepancy functions). The actual CPU time spent in the matrix multiplications and inversions were 32.59 and 11.61 seconds for the standard and improved operations respectively. Thus the ratio of the CPU time is about 2.81 as compared to the theoretically obtained ratio of 3.02.

The total CPU time spent in the execution of the programs for the standard and improved operations are 38.43 and 17.43 seconds respectively, and the balance was spent in evaluating the discrepancy functions, elements of the submatrices, and in executing the various input and output statements.

With the original discrepancy functions, proposed by Naphtali and Sandholm (refer equation 2.10), this program took 7 iterations to converge, whereas only 4 iterations were needed with the modified equation 2.2. Thus, this minor change has improved the convergence characteristics.

It was realized that the large roundoff errors in the inversion of the submatrices may impair the convergence characteristics. Three different subroutines used to study the convergence characteristics, and the results of which are as under:

	Matrix inversion subroutine	Number of iterations required
1.	Gauss Jordan with the maximum pivot-strategy	4
2.	Gauss elimination	5
3.	FO1AAF/NAG (available on DEC-10)	12

A check on the roundoff errors revealed that the FO1 AFF subroutine resulted in substantially large roundoff errors, in the matrix inversions.

It was also noted that the inversion by partitioning method resulted in less roundoff errors as compared to the standard matrix inversion, particularly in the matrices where the difference in the magnitude of the elements was vast.

Problem 2: Consider a single absorption column having 20 stages with a mixture of 4 components. (For the complete specifications please refer [1]).

This problem was solved using the conventional Thomas algorithm and a comparison of the operation counts is made in the Table 2.

A reduction in the computations by a ratio of 2.676 is obtained by sparsity exploitation, as only 17556 operations were performed instead of 46980.

This problem when implemented on DEC-10, took 4 iterations to converge to a tolerance limit of $5x10^{-8}$

TABLE 2

Comparison of Operation Counts for Problem 2

Number of components = 4

Number of stages = 20

Number of times performed	Standard operation count	Improved operation count
19	x729 = 13851	x189 = 3591
19	x729 = 13851	x 25 = 475
19	x 81 = 1539	x = 5 = 95
19	x 81 = 1539	x 45 = 855
20	x 81 = 1620	x 81 = 1620
20	x 729 = 14580	x546 = 10920
	46980	17556
	times performed 19 19 19 19 20	times operation count 19

 $\frac{\text{S.O.C.}}{\text{I.O.C.}} = 2.676$

(sum of squares of all discrepancy functions). The total CPU time spent in the matrix multiplications and inversions were 2.82 and 1.22 sec. for the standard and improved operations respectively. Thus the ratio of the CPU time is about 2.31 as against the theoretically obtained ratio of 2.676.

The total CPU time for the standard and improved programs was 7.02 and 4.34 seconds respectively, and the balance was spent in the evaluation of the discrepancy functions, the elements of the submatrices, and in executing the input and output statements.

There are a total of 19 P-blocks in the matrix after the forward substitution, each of order 9x9. Since only 19x9x5 elements of the P-blocks need to be stored instead of 19x9x9 44% saving in the storage is obtained.

Problem 3: Consider a single column with 15 stages and a 6-component mixture. In this case, as discussed in the Chapter 4, instead of specifying the heat duties of the condenser and reboiler, specifications at the 9th and 10th stages were made. The temperature of the stage 9, and the flow rate of the vapor leaving the stage 10 are specified. In the Chapter 4, two alternate approaches to solve this problem have been discussed, and the operation counts for both are compared in the Table 3.

TABLE 3

Comparison of Operation Counts for Problem 3

Number of components = 6 Number of stages = 15

	Standard	Standard operation count	unt	Improved	Improved operation count	ount
Operations	Number of times performed	Count per operation	Operation count	Number of times Op i was perfor- med	Count per Op-i	Operation count for Op i
Op 1	15	2197	32955		637	8918
. oo 3	22	2197	48334	14	85	1190
7 d0	9	2197	13182)
Op 14	4	2197	8788			:
Op 15	15	169	2535	7	<u>.</u>	180
Op 17	25	169	4225	4	. 61	1974
Op 19	15	169	2535	15	169	7535
Op 20	15	2197	32955	1.5	1562	23430
TOTAL			145509			37529

 $\frac{3.00}{1.00} = 3.877$

For the first approach which was proposed by Hofeling and Seader [2], the standard operation count is computed, for obtaining the solution by the modified Thomas algorithm using the Jacobian structumre shown in the Figure 4.

For the latter approach, the improved operation count with the sparsity exploitation was computed for obtaining the solution by the conventional Thomas algorithm. The Jacobian matrix in this case has a tridiagonal band structure.

The standard and improved operation counts are 145509 and 35729 respectively, which results in a computation reduction ratio of 3.877.

In the Hofeling-Seader approach, there are 21 P-blocks (14 tridiagonal and 7 off-tridiagonal), each of order 13x13. Whereas in the second approach there are no off-diagonal blocks and hence a total of only 14 P-blocks in Jacobian.

Therefore following the latter approach, only $1274 \ (= 14x13x7)$ elements need to be stored, whereas the standard storage requirement for the two approaches is $3549 \ (= 21x13x13)$ and $2366 \ (= 14x13x13)$ respectively.

The proposed approach is therefore more efficient, then the Hofeling-Seader approach since it requires substantially less computations and storage.

```
B<sub>1</sub> C<sub>1</sub>
                                                                                        C 1.9
A<sub>2</sub> B<sub>2</sub> C<sub>2</sub>
           A<sub>3</sub> B<sub>3</sub> C<sub>3</sub>
                      A<sub>4</sub> B<sub>4</sub> C<sub>4</sub>
                                 As Bs Cs
                                            A<sub>6</sub> B<sub>6</sub> C<sub>6</sub>
                                                        A<sub>7</sub> B<sub>7</sub> C<sub>7</sub>
                                                                   A<sub>8</sub> B<sub>8</sub> C<sub>8</sub>
                                                                              As Bs Cs
                                                                                         A10 B10 C10
                                                                                                    A_{11} \ B_{11} \ C_{11}
                                                                                                               A12 B12 C12
                                                                                                                           A<sub>13</sub> B<sub>13</sub> C<sub>13</sub>
                                                                                                                                      A<sub>14</sub> B<sub>14</sub> C<sub>14</sub>
                                                                                                                                              A<sub>15</sub> B<sub>15</sub>
                                                                                                    A_{15,10}
```

FIG 4 THE STRUCTURE OF THE JACOBIAN MATRIX (PROBLEM 3)

Problem 4: Consider the system of interlinked columns shown in the Figure 5, in which a 4-component mixture is being separated using two absorbed A₁ and A₂ having 10 plates each, and two distillation columns having 15 and 12 plates respectively. The arrangement of columns and ordering of the trays is also shown in the same figure, and the structure of the Jacobian matrix is presented in the Figure 6. (For the complete specifications please refer Ketchum [3]).

The modified Thomas algorithm was employed to solve the system, and the various steps of which are presented in the Chapter 2.

By exploiting the sparsity a reduction in the computations by a ratio of 3.013 was obtained, as only 62961 operations were performed instead of 189702 standard operations. The comparison and details of the standard and improved operation counts have been presented in the Table 4.

There are 46 upper diagonal, 53 off-tridiagonal, and hence a total of 99 P-blocks in the matrix, after the forward substitution. Each P-block is of the order 9x9. As discussed earlier only a 9x5 matrix for every P-block needs storing, since the remaining elements are zero, and hence a 44% saving in the storage may be realized.

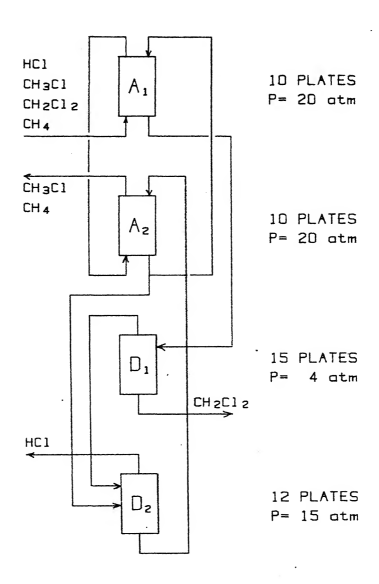


FIG 5 THE ARRANGEMENT OF THE INTERLINKED-COLUMNS (PROBLEM 4)

```
B<sub>1</sub> C<sub>1</sub>
                                                                                              C 120
 A₂ B₂ C₂
     A<sub>3</sub> B<sub>3</sub> C<sub>3</sub>
          A, B, C,
                As Bs Cs
                    A6 B6 C6
                         A7 B C7
                              As Bs Cs
                                   Ag Bg Cg
                                        A10 B10
                                                  B11 C11
                                                                                                                                                                                                                             C 11.47
                                                  \Lambda_{12}\,B_{12}\,C_{12}
                                                      A<sub>13</sub> B<sub>13</sub> C<sub>13</sub>
                                                           A14 B14 C14
                                                                A<sub>15</sub> B<sub>15</sub> C<sub>15</sub>
                                                                     A16 B16 C16
                                                                          A17 B17 C17
                                                                              A18 B18 C18
                                                                                   A19 B19 C19
A zali
                                                                                        A20 B20
                                                                                                  B<sub>21</sub> C<sub>21</sub>
                                                                                                  Azz Bzz Czz
                                            A 23,10
                                                                                                       A23 B23 C23
                                                                                                          A24 B24 C24
                                                                                                               Az Bz Cz
                                                                                                                    A26 B26 C26
                                                                                                                         Azı Ber Czı
                                                                                                                              Aze Bze Cze
                                                                                                                                   Az9 Bz9 Cz9
                                                                                                                                        A30 B30 C30
                                                                                                                                             A31 B31 C31
                                                                                                                                                  A32 B32 C32
                                                                                                                                                       A33 B33 C33
                                                                                                                                                            A34 B34 C34
                                                                                                                                                                 A35 B35
                                                                                                                                                                          B<sub>36</sub> C<sub>36</sub>
                                                                                                                                                                          Азт Взт Сэт
                                                                                                                                                                               A38 B38 C38
                                                                                                                                                                                    A39 B39 C39
                                                                                           A 40,2i
                                                                                                                                                                                         A40 B40 C40
                                                                                                                                                                                              A4; B41 C41
                                                                                                A 41,20
                                                                                                                                                                                                   A42 B42 C42
                                                                                                                                                                                                       A43 B43 C43
                                                                                                                                                                                                            A44 B44 C44
                                                                                                                                                                                                                  A45 B45 C45
                                                                                                                                                                                                                       A46 B46 C46
                                                                                                                                                                                                                           A47 B47
```

FIG 6 THE STRUCTURE OF THE JACOBIAN MATRIX FOR PROBLEM 4

TABLE 4

Comparison of Operation Counts for Problem 4

Number of components = 4

Total number of stages = 47

Operations	Number of times performed		Standard operation count	Improved operation count
Op 1	42	x729	= 30618	x225 = 9450
Op 2	2	x729	= 1458	x 81 = 162
0p 3	43	x729	= 31347	x 41 = 1763
0p 4	38	x729	= 27702	x = 41 = 1558
0p 6	2	x729	= 1458	x125 = 250
Op 8	36	x729	= 26244	x225 = 8100
Op 10	1	x729	= 729	x405 = 405
Op 11	25	x729	= 18225	x225 = 5625
Op 12	1 *	x729	= 729	x225 = 225
Op 15	45	x 81	= 3645	x = 9 = 405
Op 16	2	x 81	= 162	x 25 = 50
Op 17	66	x 81	= 5346	x 45 = 2970
0p 18	40	x 81	= 3240	x 45 = 1800
Op 19	47	x 81	= 3807	x 81 = 3807
Op 20	47	x729	= 34263	x546 = 25662
Standard matrix multiplica- tion	1	x729	= 729	x729 = 729
Total	ng at the first of the state of	and the second s	189702	62961

 $\frac{\text{S.O.C.}}{\text{I.O.C.}} = 3.013$

It may be noted here that out of the 99 P-blocks, 15 are null matrices ($P_{11,20}$ through $P_{18,20}$, $P_{21,47}$, $P_{22,47}$, and $P_{35,47}$ through $P_{39,47}$) and hence need not be stored, thereby leaving only 85 P-blocks which must be stored. Therefore, only 3825 (= 85x9x5) elements are stored instead of 8019 (= 99x9x9), and a 52% of actual saving in the storage is obtained.

Problem 5: Consider the system of interlinked columns for separating a mixture of 3-components using 3 columns which are connected as shown in the Figure 7. These columns are assumed to be split into section 2a having 4 plates, 2b having 3 plates, 3a having 4 plates, 1 having 4 plates, and 3b having 4 plates, and are rearranged in that order. The Jacobian matrix for this system is presented in the Figure 8. The modified Thomas algorithm was applied to obtain the solution.

In this case, the sparsity exploitation has resulted in a reduction in the computations by a ratio of 2.744, as only 13393 operations were performed instead of the 36750. The comparison and details of the operation counts are presented in the Table 5.

There are 18 upper diagonal, 11 off-tridiagonal, and hence a total of 29 P-blocks, each of order 7x7, in the Jacobian matrix, (P_1 through P_{18} , $P_{4,12}$ through $P_{4,10}$, and $P_{11,16}$ through $P_{14,16}$) as shown in the Figure 5. Since only

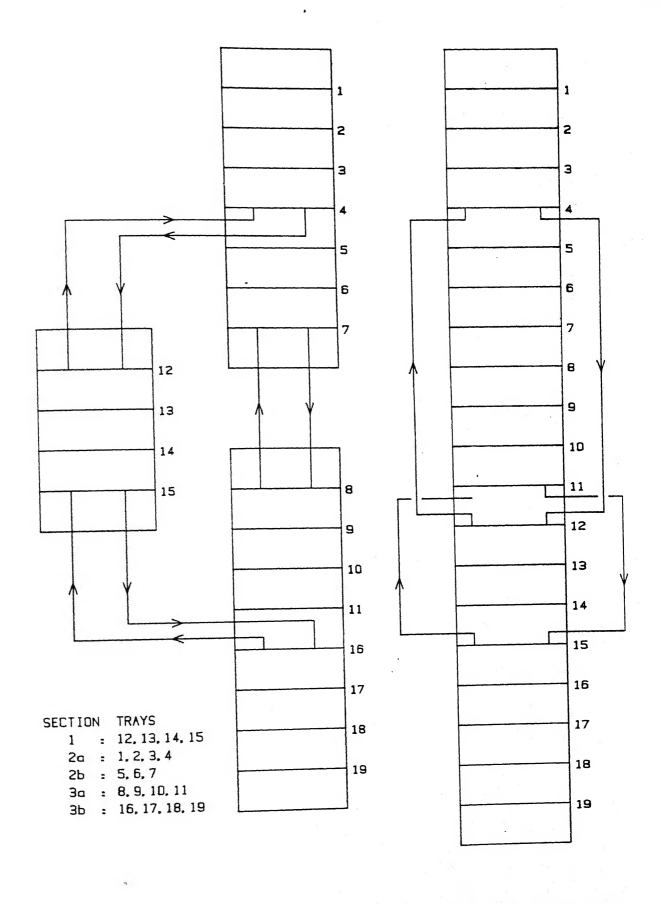


FIG 7 AN ARRANGEMENT OF INTERLINKED-COLUMNS (PROBLEM 5 AND 6)

```
B_1 C_1
Az Bz Cz
           A<sub>3</sub> B<sub>3</sub> C<sub>3</sub>
                      A<sub>4</sub> B<sub>4</sub> C<sub>4</sub>
                                                                                                                        C 4,12
                                 A<sub>5</sub> B<sub>5</sub> C<sub>5</sub>
                                            As Bs Cs
                                                       A<sub>7</sub> B<sub>7</sub> C<sub>7</sub>
                                                                  A<sub>e</sub> B<sub>e</sub> C<sub>e</sub>
                                                                             Ag Bg Cg
                                                                                        A<sub>10</sub> B<sub>10</sub> C<sub>10</sub>
                                                                                                  A<sub>11</sub> B<sub>11</sub>
                                                                                                                                                                     C 11,16
                                A 12,4
                                                                                                                      B12 C12
                                                                                                                         A<sub>13</sub> B<sub>13</sub> C<sub>13</sub>
                                                                                                                                    A<sub>14</sub> B<sub>14</sub> C<sub>14</sub>
                                                                                                                                               A<sub>15</sub> B<sub>15</sub> C<sub>15</sub>
                                                                                                              A 16,11
                                                                                                                                                          A<sub>16</sub> B<sub>16</sub> C<sub>16</sub>
                                                                                                                                                                     A<sub>17</sub> B<sub>17</sub> C<sub>17</sub>
                                                                                                                                                                                 A<sub>18</sub> B<sub>18</sub> C<sub>18</sub>
                                                                                                                                                                                            A19 B19
```

FIG 8 THE STRUCTURE OF THE JACOBIAN MATRIX (PROBLEM 5 AND 6)

TABLE 5

Comparison of Operation Counts for Problem 5

Number of components = 3

Total Number of stages = 19

Operation ,	Number of times performed	Standard operation count	Improved operation count
Op 1	18	x 343 = 6174	x112 = 2016
Op 3	30	x343 = 10290	x25 = 750
Op 7	10	x343 = 3430	x112 = 1120
Op 9	1	x343 = 343	x196 ≠ 196
Op 14	18	x343 = 6174	x112 = 2016
Op 15	19	x 49 = 931	x 7 = 133
Op 17	40	x 49 = 1 960	x 28 = 1120
Op 19	19	x 49 = 931	x 49 = 931
Op 20	19	x343 = 6517	x269 = 5111
Total		36750	13393

 $\frac{5.0.C.}{1.0.C.} = 2.744$

a total of 812 (= 29 x 7 x 4) elements are stored instead of the 1421 (= 29x7x7), a saving of about 43% in the storage requirement is obtained.

Problem 6: The same system of the interlinked columns described in the problem 2 was solved with a 6-component mixture. The structure of the Jacobian and the method of solution remains the same, but the order of each submatrix in the Jacobian is 13x13 now.

In this case, the sparsity exploitation has resulted in a reduction in the computations by a ratio of 3.21, which is more significant than that in the problem 2. Only 69811 operations were performed instead of the 224094, and the details are presented in the Table 6.

The total number of P-blocks is 29 (same as in problem 2). A total of only 2693 (= 29x13x7) elements are stored instead of 4901 (= 29x13x7), and thereby a saving of 46% can be realized in the storage requirements.

TABLE 6

Comparison of the Operation Counts for Problem 6

Number of components = 6

Total number of stages = 19

Operation	Number of times performed	Standard operation count	Improved operation count
Op 1	18	x2197 = 39546	x637 = 11466
Op 3	30	x2197 = 65910	x 85 = 2550
Op 7	10	x2197 = 21970	x637 = 6370
Op 9	1	x2197 = 2197	x1183= 1183
Op 14	18	x2197 = 39546	x637 = 11466
Op 15	19	x 169 = 3211	x 13 = 247
Op 17	40	x 169 = 6760	x 91 = 3640
Op 19	19	x 169 = 3211	x169 = 3211
Op 20	19	x2197 = 41743	x1562= 29678
COLUMN THE STATE OF THE STATE O	gadinin min ang ang ang ang ang ang ang ang ang an	224094	69811
kasaliguselija, salinsistelekselikselikselikselikselikselikseliks	and the second seco	S.O.C. = 3.21	устандын түү төрсөн байгай Ангору и Эмен Айгий Жий Айра айры айри айрий байгийн айгийн айгийн айгийн айгийн а

CHAPTER 7

CONCLUSIONS

In the algorithm presented in this work the sparsity and the structure of the submatrices was exploited in the various matrix multiplications and inversions, and thereby a significant reduction in the operation-count and storage requirements was obtained. The reduction in the computations becomes increasingly significant with the increase in the number of components.

Theoretically, a maximum of 3.43 (= 24/7) times reduction in the computations may be realized in the single column problems if the number of components is very large. In general, a reduction in storage requirement by a ratio of (c+1)/(2c+1) was also realized. The exploitation of sparsity in solving the single column problems by the conventional block Thomas algorithm has shown 2.676 and 3.02 times improvement in the operation-count, in the test problems 1 and 2.

An efficient approach to solve the problems with the intermediate tray specifications has been proposed and shown to be more advantageous than the one proposed by Hofeling and Seader [2], in both the computational and the storage aspects. For instance, in the problem 3, the proposed approach has been proved to be 3.877 times computationally efficient and 64% more storage-saving than

the Hofeling-Seader approach.

In the solution of a system of interlinked columns by the modified Thomas algorithm, the exploitation of sparsity has resulted in a 3.013, 2.744 and 3.21 times improvement in the operation count, in the problems 4,5 and 6 respectively. Though this ratio varies from problem to problem, and improves with the increase in the number of components, in general it is close to 3.

The saving, at least by a ratio of (c+1)/(2c+1) in the storage requirement is also realized in the systems of interlinked columns, which significantly improves further if some of the off-tridiagonal P-blocks are null matrices.

The proposed method of the sparsity exploitation can be applied to the other related techniques, for instance, the convergence domain extension methods proposed by Vickery and Taylor [8]; and the other methods of solution proposed by Kubicek [7], and Stadtherr [5].

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APPENDIX

ELEMENTS OF THE SUBMATRICES

TRIDIAGONAL SUBMATRICES:

(i) For Matrix A

$$\frac{\partial M_{j,i}}{\partial I_{j-1,k}} = r_{Lj-1} \delta_{ik}; \quad \frac{\partial M_{j,i}}{\partial V_{j-1,k}} = 0; \quad \frac{\partial M_{j,i}}{\partial T_{j-1}} = 0$$

$$\frac{\partial Q_{j,i}}{\partial I_{j-1,k}} = \frac{\partial Q_{j,i}}{\partial V_{j-1,k}} = \frac{\partial Q_{j,i}}{\partial T_{j-1}} = 0$$

$$\frac{\partial E_{j}}{\partial T_{j-1,i}} = r_{Lj-1}h_{j-1,i}; \frac{\partial E_{j}}{\partial V_{j-1,i}} = 0; \frac{\partial E_{j}}{\partial T_{j-1}} = r_{Lj-1}$$

$$c$$

$$\sum_{i=1}^{C} t_{j-1,i} \frac{\partial h_{j-1,i}}{\partial T_{j-1}}$$

(ii) For Matrix B

$$\frac{\partial M_{j,i}}{\partial T_{j,k}} = -(1+s_j)\delta_{i,k}; \quad \frac{\partial M_{j,i}}{\partial V_{j,k}} = -(1+s_j)\delta_{i,k}; \quad \frac{\partial M_{j,i}}{\partial T_{j}} = 0$$

$$\frac{\partial Q_{j,i}}{\partial l_{j,k}} = \frac{\eta_{j}}{L_{j}^{2}} \begin{bmatrix} K_{j,i} \delta_{i,k} L_{j} + l_{j,i} & \sum_{p=1}^{c} \left(\frac{\partial K_{i,i}}{\partial x_{j,p}} \right) \\ (\delta_{k,p} - \frac{l_{j,p}}{L_{j}}) - K_{j,i} l_{j,i} \end{bmatrix}$$

$$\frac{\partial Q_{j,i}}{\partial V_{j,k}} = \frac{1}{V_{j}} \begin{bmatrix} \frac{\eta_{j} l_{j,i}}{L_{j}} & \sum_{p=1}^{c} (\frac{\partial K_{j,i}}{\partial Y_{j,p}})(\delta_{p,k} - \frac{V_{j,p}}{V_{j}}) \\ -\delta_{i,k} + \frac{V_{j,i}}{V_{j}} \end{bmatrix}$$

$$\frac{\partial Q_{j,i}}{\partial T_{j}} = \frac{\eta_{j} l_{j,i}}{L_{j}} \left(\frac{\partial K_{j,i}}{\partial T_{j}}\right)$$

$$\frac{\partial E_{j}}{\partial l_{j,i}} = -(1 + s_{j})h_{j,i}; \frac{\partial E_{j}}{\partial v_{j,i}} = -(1 + S_{j})H_{j,i}$$

$$\frac{\partial E_{j}}{\partial T_{j}} = -(1 + s_{j}) \sum_{i=1}^{C} 1_{j,i} \frac{\partial h_{j,i}}{\partial T_{j}} - (1 + S_{j}) \sum_{i=1}^{C} v_{j,i} \frac{\partial H_{j,i}}{\partial T_{j}}$$

(iii) For Matrix C

$$\frac{\partial M_{j,i}}{\partial I_{j+1,k}} = 0; \quad \frac{\partial M_{j,i}}{\partial V_{j+1,k}} = r_{Vj+1} \hat{\delta}_{i,k}; \quad \frac{\partial M_{j,i}}{\partial T_{j+1}} = 0$$

$$\frac{\partial Q_{j,i}}{\partial I_{j+1,k}} = 0; \quad \frac{\partial Q_{j,i}}{\partial T_{j+1}} = 0$$

$$\frac{\partial Q_{j,i}}{\partial v_{j+1,k}} = \frac{(1-\eta_j)r_{Vj+1}}{D} \left[\delta_{i,k} - \frac{r_{Vj+1}v_{j+1,i}+R_{i,v}v_{V,i}+R_{j,v}v_{Zi}}{D}\right]$$

$$D = r_{Vj+1} V_{j+1} + R_{jy} V_{y} + R_{jz} V_{z}$$

$$\frac{\partial E_{j}}{\partial I_{j+1,k}} = 0; \quad \frac{\partial E_{j}}{\partial V_{j+1,i}} = r_{Vj+1} H_{j+1,i}$$

$$\frac{\partial E_{j}}{\partial T_{j+1}} = r_{Vj+1} \sum_{i=1}^{z} v_{j+1,i} \left(\frac{\partial H_{j+1,i}}{\partial T_{j+1}} \right)$$

OFFDIAGONAL SUBMATRICES:

(i) For A-matrix:

$$\frac{\partial M_{j,i}}{\partial l_{p,k}} = R_{j,p} \delta_{i,k}; \frac{\partial M_{j,i}}{\partial V_{p,k}} = \frac{\partial M_{j,i}}{\partial T_{p}} = 0$$

$$\frac{\partial Q_{j,i}}{\partial I_{p,k}} = \frac{\partial Q_{j,i}}{\partial V_{p,k}} = \frac{\partial Q_{j,i}}{\partial T_{p}} = 0$$

$$\frac{\partial E}{\partial I_{p,k}} = R_{j,p}h_{p,i}; \quad \frac{\partial E_{j}}{\partial V_{p,k}} = 0; \quad \frac{\partial E_{j}}{\partial T_{p}} = R_{j,p} \quad \frac{\Sigma}{i=1} \quad \frac{\partial h_{p,i}}{\partial T_{p}}.$$

(ii) For C-matrix:

$$\frac{\partial M_{j,i}}{\partial I_{y,k}} = 0; \quad \frac{\partial M_{j,i}}{\partial V_{y,k}} = R_{j,y} \delta_{i,k}; \quad \frac{\partial M_{j,i}}{\partial T_{y}} = 0$$

$$\frac{\partial Q_{j,i}}{\partial I_{y,k}} = \frac{\partial Q_{j,i}}{\partial T_{y}} = 0$$

$$\frac{\partial Q_{j,i}}{\partial V_{j,k}} = \frac{(1-\eta_j)R_{j,Y}}{D} \left[\delta_{i,k} - \frac{r_{Vj+1}V_{j+1,i} + R_{j,Y}V_{Y,i} + R_{j,Z}V_{Z,i}}{D}\right]$$

where

$$D = r_{Vj+1} V_{j+1} + R_{j,y} v_{y,i} + R_{j,z} v_{z,i}$$

$$\frac{\partial E_{j}}{\partial I_{y,i}} = 0; \quad \frac{\partial E_{j}}{\partial v_{y,i}} = R_{j,y} H_{y,i}; \quad \frac{\partial E_{j}}{\partial T_{y}} = R_{j,y} \sum_{i=1}^{c} v_{y,i}$$

$$(\frac{\partial H_{y,i}}{\partial T_{y}}).$$

APPENDIX B1 (SAMPLE PROBLEM NO.1)

K values for the temperature range of - 25°F to 40°F and at a pressure of 800 lb/in²abs

Component	811	B21	a ₃₁	a ₄₁
17日本・日本の日本の日本の日本の日本の日本の日本の日本の日本の日本の日本の日本の日本の日	- 0.6282223x10 ⁻¹	0.30688802x10 ⁻³	0.39996468x10 ⁻⁶	-0.57899830x10 ⁻⁹
	0.50596821	. 1	-0.15009991x10 ⁻⁵	0.34494154x10 ⁻⁸
	0.15584934	-0.15205775x10 ⁻³	0.50349212x#0 ⁻⁶	-0.17713546x10 ⁻⁹
O.H.	0.91486037x10 ⁻¹		0.33741924x10 ⁻⁶	0.14797150x10-9
0 7 0 °C	0.37769508x10-1		0.29233627x10-6	-0.48597680x10-11
10,H,	0.36708355x10 ⁻¹		0.28026648x10 ⁻⁶	0.10462797x10 ⁻¹⁰
n C, H,	0.37231278x10 ⁻¹		0.37584653x10 ⁻⁶	-0.69237741x10 ⁻¹⁰
4 - C	0.15414596x10.1		0.12591028x10 ⁻⁶	0.73157133x10-10
1 C.H.	0.19747034x10.1		0.14439195x10 ⁻⁶	0.56656790x10-10
n C.H.	0.88765752x10 ⁻³		-0.40746951x10 ⁻⁷	0.15187203x10-9
n C.H.	0.63677356x10 ⁻²	1	$0.31795974x10^{-7}$	0.78284379x10-10
D O H.	0.99674799x10 ⁻²		0.82305291x10-7	0.21022392x10-10
ಕ್ಷಗತ್ತಿ ಗೆಗ್ಗೆ	0.78793392x10-2		$0.62435951x10^{-7}$	0.25793478x10-10
n C. H.	0.64146556x10 ⁻²		0.30005250x10 ⁻⁷	0.30266026x10-10

 $K_1 = T (a_{11} + a_{21} T + a_{31} T^2 + a_{41} T^3)^3 (T in ^R)$

LIGHTD ENTHALPIES FOR THE TEMPERATURE RANGE OF - 25°F to 40°F at P = 800 lb/in abs. Appendix Bl continued.

Andrew Community and the community of th		کو	ع	ŗ
Component	D11	⁰ 21	"31	741
009	0.22524075x10 ⁴	0.5446243x10	$0.2791080x10^{-1}$	- 0.18765335x10 ⁻⁴
N	0.15837112x104	0.3731512x10 ¹	0.17655857x10 ⁻¹	- 0.14662071x\$0-4
CH,	0.81635181x10 ³	0.7206460x10 ¹	0.15354034x10 ⁻¹	- 0.84406456x10 ⁻⁵
o, He	0.97404712x10 ³	0.11454294x10 ²	0.79399594×10 ⁻²	- 0.42183183x10 ⁻⁶
O.H.	$0.21237510x10^4$	0.46583524x101	0.31726830x10 ⁻¹ .	$-0.12580301x10^{-4}$
10,H40	0.17543628x104	0.92456856x10 ¹	0.30206113x10 ⁻¹	- 0.89584664x10 ⁻⁵
n O, H, O	0.32309192x104	0.66175545x10 ¹	0.38262386x10 ⁻¹	- 0.16110935x10 ⁻⁴
F C. H.	0.33611663x104	0.39552670x10 ¹	0.54925641x10 ⁻¹	- 0.25869682x10 ⁻⁴
n C _E H ₄₀	0.43454375x104	0.10596339x10 ²	0.43731511x10-1	- 0.19637475x10 ⁻⁴
n Och,	-0.44150469x104	0.70354599x10 ²	-0.67470074x10 ⁻¹	0.60245657x10 ⁻⁴
n C, H,c	0.66707016x10 ²	0.18159073x10 ²	0.38164884x10 ⁻¹	- 0.42837073x10 ⁻⁵
B C _R H ₄ R	-0.10632578x10 ²	0.19229950x10 ²	0.40186413x10 ⁻¹	- 0.70521889x10 ⁻⁶
B CaH2O	-0.79141992x104	$0.81615143x10^{2}$	-0.79501927x10 ⁻¹	0.83943509x10-4
n 040H22	-0.67810352x10 ⁴	0.74108551x10 ²	-0.58315706x10 ⁻¹	0.75087155x10-4

h₁ = b₁₁ + b₂₁T + b₃₁T² + b₄₁T³ (Tin °R) Btu/lb mole.

To 40° F AT P = 800 lb/in²abs. Appendix Bicontinued. VAPOR ENTWILDIES FOR THE TEMPERATURE RANGE OF - 25°F

p apparatuate abustinistimitimitis despe	₹_	77	17	41
°00	0.13978977×10 ⁵	$-0.96359463x10^{1}$	0.38228422x10 ⁻¹	-0.26870170x10 ⁻⁴
	0.48638672x104	$-0.21227379x10^{1}$	0.17565668x10 ⁻¹	-0.11367006x1c ⁻⁴
	0.63255430x104	- 0.20747757x101	0.18532634x10-1	-0.10630416x10 ⁻⁴
	0.10628934x10 ⁵	- 0.28718834x10 ¹	0.24877094x10 ⁻¹	-0.13233222x10-4
OZHO.	0.13954383x10 ⁵	- 0.41930256x10 ¹	0.32614145x10 ⁻¹	-0.15483340x10 ⁻⁴
10 H	0.94088984x104	0.39262680x10 ²	-0.55596594x10 ⁻¹	●-51507392×10 ⁻⁴
n C.H.O.	0.57302344x104	0.75117737x10 ²	-0.13120884x10 ⁰	0.10517908x10 ⁻³
1 C.H.	0.83081953x104	0.75267792x10 ²	_0.12945843x10 ⁰	0.10845697x10 ⁻³
D C L	0.12804211x105	0.61654007x10 ²	-0.97365201x10 ⁻¹	0.84398722x10-4
n C H,	0.23001684x10 ⁵	$0.27744919x10^2$	-0.31545494x10 ⁻¹	0.49981289x10-4
n G _r H _r C	0.14876816x10 ⁵	0.59342438x10 ²	-0.81853271x10 ⁻¹	0.81429855x10-4
n C _o H ₂	0.32793215x10 ⁵	$-0.35040283x10^{2}$	0.11162955x10 ⁰	-0.42647429x10 ⁻⁴
n CoHoO	0.47024656x10 ⁵	- 0.95395035x10 ²	0.24547529x10 ⁰	-0.13209638x10 ⁻³
n C ₁₀ H ₂₂	0.55238211x10 ⁵	- 0.13195618x10 ³	0.32518369x10 ⁰	-0.18188384x10 ⁻³

(T in oR) Btu/lb mole. H1 = C11 + C21 T + C31 T + C41 T

APPENDIX B2

SAMPLE PROBLEM No.2

DA DA

K values

Material	Temperatu	re
	100°F	200°F
A	500.0	55 0. 0.
B	1.50	1.8
C	0.90	1.00
D	1.0×10^{-6}	1.5 x 10
	Molar liquid enthalpi	es, 10 ³ cal/mol
A	0.01	0.013
В	0.30	0.33
C	0,40	0.44
D	1.50	1.90
	Molar vapor enthalpie	es, 10 ³ cal/mole
A	1.00	1.002
В	1.80	1.82
C	2,00	2.03

```
INTERLINKED COLUMNS
                                                     PROGRAM INTLNK
INTEGER C,CP1,TWOC,CT
DIMENSION JCOND(2),1TYPEC(2),JREBL(2),ITYPER(2),
1 IX(6),IY(6),LORY(6),RATIO(6),JNOSY(3),JNOSL(3),
1 JVAPF(1),TFV(1),FEEDV(1,1),JLIQF(1),TFL(1),FEEDL(1,1),
1 JSPC(4),LBL(4),SPCVAL(4),ICOMP(4),
1 QG(12),SV(12,2),Sb(12,2),SSV(12),SSL(12),T(12),ETA(12),
1 HV(4),HL(4),AK(4),VECTOR(4),
1 BV(47),BL(47),RV(47),RL(47),DENOM(47),
DIMENSION A(9,9),B(9,9),CC(9,9),AOFF(9,9),COFF(9,9),DIMENSION ALFA(9,5),BETA2(9,5),BETA1(9),VEC1(9),DIMENSION F(47,9),P(85,9,5),Q(47,9)
                                                   DIMENSION F(47;9),P(85,9,5),Q(47;9)

OPEN(UNIT=33,DEVICE='DSK',FILE='INP.IN')

OPEN(UNIT=51;DEVICE='DSK',FILE='TH.OUT')

READ(33,*)N

READ(33,*)NOF,NABOVE

READ(33,*)NCOND,NREBL,NUV,NOL,NVAPF,NLIQF,NSPC

IF(NCOND_LT.1)GOTO 810

DO-800 I=1,NCOND

READ(33,*)JCOND(I),ITYPEC(I)

IF(NREBL.LT.1)GOTO 812

DO-801 I=1,NREBL

READ(33,*)JREBL(I),ITYPER(I)

IF(NSPC.LT.1)GOTO 812

DO-802 I=1,NSPC

READ(33,*)JNEBL(I),LBL(I),SPCVAL(I),ICOMP(I)

IF(NOV.LT.1)GOTO 813

READ(33,*)JNOSV(I),I=1,NOV)

IF(NOL.LT.1)GOTO 814

READ(33,*)(JNOSV(I),I=1,NOL)

IF(NOFF.LT.1)GOTO 815

DO-803 I=1,NOFF

READ(33,*)JXAPF(I),TFV(I),(FEEDV(I,KK),KK=1,C)

LF(NEIGF.LT.1)GOTO 817

DO-805 I=1,NLIQF

READ(33,*)JLIQF(I),TFL(I),(FEEDL(I,KK),KK=1,C)

CONTINUE

READ(33,*)(GQ(J),J=1,N)

READ(33,*)(SSV(J),J=1,N)
 800
 810
 801
811
802
812
 813
  814
  803
  815
   804
   816
   805
   817
                                                         CONTINUE
READ(33,*)(QQ(J),J=1,N)
READ(33,*)(SSV(J),J=1,N)
READ(33,*)(SSL(J),J=1,N)
READ(33,*)(ETA(J),J=1,N)
READ(33,*)(T(J),J=1,N)
READ(33,*)((SV(J,I),I=1,C),J=1,N)
READ(33,*)((SL(J,I),I=1,C),J=1,N)
                                                            ICHK=0
KCOND=0
                                                           KCOND=0
KREBL=0
NTOT=NCOND+NREBL
DO 110 KK=1,NSPC
IF(LBL(KK).LT.3)GOTO 110
J=JSPC(KK)
DO 111 MM=1,NCOND
IF(J.EQ.JCOND(MM))GOTO 113
CONTINUE
DO 112 MM=1,NREBL
      111
                                                            DO 112 MM=1, NREBL
IF(J.EQ.JREBL(MM))GOTO 114
CONTINUE
GOTO 115
KCOND=KCOND+1
      112
      113
```

```
GOTO 115
                               KREBL=KREBL+1
ICHK=ICHK+1
114
115
110
                               CONTINUE
                               ISTP=0
                                IF (KCOND. EQ. NCOND) GOTO 116
                               ISTP=1
TYPE 901, KCOND, NCUND
IF(KREBL.EQ.NREBL)GOTO 117
ISTP=1
116
                               TYPE 902, KREBL, NREBL
IF(ICHK.EQ.NTOT)GUTO 119
IF(ICHK.LT.NTOT)GUTO 118
TYPE 904, ICHK, NTOT
117
                               STOP
TYPE 903, ICHK, NTOT
118
                               STOP
                               FORMAT(1X, ---ERRUR---/1X, INFORMATION ABOUT ',12, CONDENSERS'

1, "HAS BEEN PROVIDED WHEREAS THERE ARE ",12, "CONDENSERS'

FORMAT(1X, ---ERRUR---'/1X, INFORMATION ABOUT ',12, "REBOILERS'

1, "HAS BEEN PROVIDED WHEREAS THERE ARE ",12, "REBOILERS'

FORMAT(1X, ---ERRUR---'/1X, NO. OF SPECIFICATIONS SHOULD BE ',12

1, "BUT ",12," HAVE BEEN PROVIDED '/1X, "PROBLEM UNDERSPECIFIED')

FORMAT(1X, ---ERRUR---'/1X, NO. OF SPECIFICATIONS SHOULD BE ',12

1, "BUT ",12," HAVE BEEN PROVIDED '/1X, "PROBLEM OVERSPECIFIED')

IF(ISTP-EU,1)STUP
901
902
903
904
119
                               DO 145 J=1,N
RL(J)=1.
RV(J)=1.
DO 146 KK=1,NOFF
145
                               J=IY(KK)

IF(LORV(KK).EQ.0)GOTO 147

RV(J)=RV(J)=RATIO(KK)

GOTO 146

RL(J)=RL(J)=RATIO(KK)

CONTINUE

DO 151 J=1, N

BV(J)=0
147
                               DU 131 0-1,N
BV(J)=0.
BL(J)=0.
DU 151 I=1,C
BV(J)=BV(J)+SV(J,I)
BL(J)=BL(J)+SL(J,I)
DU 200 J=1,N
151
                               JM1=J-1
                                JP1=J+1
                               JP1=J+1
TJ=T(J)
SD0==(1.+SSL(J))
SD1==(1.+SSV(J))
DO 152 KK=1,NCOND
IF(J.EQ.JCOND(KK))GUTO 180
CONTINUE
152
                                CUNTINGE

DO 153 KK=1,NREBL

IF(J:EQ.JREBL(KK))GOTO 190

CONTINUE

CALL ENV(TJ,HV)

CALL ENL(TJ,HL)

CALL FINDK(TJ,AK)

SUMV=0.
 153
                               SUMV=0.

SUML=0.

COEF=ETA(J)*BV(J)/BL(J)

DO 154 I=1,C

F(J,I)=SD0*SL(J,I)+SD1*SV(J,I)

F(J,I+C)=CUEF*AK(1)*SL(J,I)*SV(J,I)

SUMV=SUMV+HV(I)*SV(J,I)

SUML=SUML+HL(I)*SL(J,I)

F(J,CT)=SD0*SUML+SD1*SUMV+QQ(J)

IF(J.EQ.1)GUTO 170
 154
```

```
3
```

```
DO 155 KK=1,NOL
IF(J.EQ.JNOSL(KK))GOTO 170
CONTINUE
155
                   CALL ENL(T(J-1), HL)
SUMHL=0.0
DO 157 I=1, C
F(J,I)=F(J,I)+SL(JM1,I)*RL(JM1)
SUMHL=SUMHL+HL(I)*SL(JM1,I)
F(J,CT)=F(J,CT)+SUMHL*RL(JM1)
IF(J,EQ,N)GOTO 200
DO 171 KK=1,NOV
IF(J,EQ,JNOSV(KK))GOTO 200
CONTINUE
CALL ENV(T(J+1),HV)
SUMHV=0.
                    CALL ENL(T(J-1), HL)
157
170
171
                    SUMHV=0.

DO 173 I=1,C

F(J,1)=F(J,I)+SV(JP1,I)*RV(JP1)

SUMHV=SUMHV+HV(I)*SV(JP1,I)

F(J,CT)=F(J,CT)+SUMHV*RV(JP1)

GOTO 200
173
CONDENSER PART
                   SER PART
DO 181 I=1,C
F(J,I)=SV(JP1,I)*RV(JP1)+SD1*SV(J,I)+SD0*SL(J,I)
IF(ITYPEC(KK).EQ.0)GOTO 182
DO 184 I=1,C
F(J,I+C)=BV(J)*SL(J,I)/BL(J)-SV(J,I)
GOTO 200
CALL FINDK(TJ,AK)
COEF1=ETA(J)*BV(J)/BL(J)
COEF2=(1.-ETA(J))*BV(J)/BV(JP1)
DO 183 I=1,C
F(J,I+C)=COEF1*AK(I)*SL(J,I)-SV(J,I)+COEF2*SV(JP1,I)
GOTO 200
181
184
182
183
FEED PART OF THE DISCREPENCY FUNCTIONS
                     DO 201 KK=1, NLIQF
                      J=JLIOF(KK)
                     TF=TFL(KK)

CALL ENL(TF, HL)

DO 201 I=1, C

F(J, I)=F(J, I)+FEEDL(KK, I)

F(J, CT)=F(J, CT)+HL(I)*FEEDL(KK, I)

DO 202 KK=1, NVAPF

J=JVAPF(KK)
 201
                      J=JVAPF(KK)
TF=TFV(KK)
CALL ENV(TF,HV)
DD 202 1=1,C
F(J,I)=F(J,I)+FEEDV(KK,I)
F(J,CT)=F(J,CT)+HV(I)*FEEDV(KK,I)
 202
     CONTRIBUTION OF INTERLINKS TO THE DISCREPENCY FUNCTIONS
                      DO 210 KK=1, NOFF
```

```
J=1X(KK)
JK=1Y(KK)
TK=T(JK)
IF(LORV(KK).EQ.0)GOTO 211
CALL ENV(TK,HV)
SUMV=0.
DO 213 1=1,C
F(J,1)=F(J,1)+SV(JK,1)*RATIO(KK)
SUMV=SUMV+HV(I)*SV(JK,I)
F(J,CT)=F(J,CT)+SUMV*RATIO(KK)
GOTO 210
CALL ENL(TK,HL)
SUML=0.
213
211
                            SUML=0.
DO 212 I=1,C
F(J,I)=F(J,I)+SL(JK,I)*KATIO(KK)
SUML=SUML+SL(JK,I)*HL(I)
F(J,CT)=F(J,CT)+SUML*RATIO(KK)
CONTINUE
212
210
    LAST TERM OF THE EQUILIBRIUM DISCREPENCY FUNCTION
                           DO 240 J=1,N
IF(ETA(J).EQ.1.)GOTO 240
DO 220 KK=1,NCOND
IF(J.EQ.JCOND(KK))GUTO 240
CONTINUE
DO 221 KK=1,NREBL
IF(J.EQ.JREBL(KK))GUTO 240
CONTINUE
KK=1
220
221
                             KK=1
                             IF(J.NE.IX(KK))GOTO 223
IF(LORV(KK).EG.1)GOTO 230
222
                            KK=KK+1

IF(KK.LE.NOFF)GOTO 222

DO 225 KK=1,NOV

IF(J.EQ.JNOSV(KK))GOTO 226

CONTINUE
223
225
                            CONTINUE
COEFF=(1.-ETA(J))*BV(J)/BV(J+1)
DO 227 I=1,C
F(J,I+C)=F(J,I+C)+CUEFF*SV(J+1,I)
DENUM(J)=RV(J+1)*BV(J+1)
GOTO 240
TYPE 905,J
FORMAT(1X,'--ERRUR---'/1X,'STAGE ',I2,' DOES NOT HAVE ANY
1 'VAPOR INPUT AND HAS NOT BEEN DEFINED AS A REBOILER')
STOP
227
226
905
                              STOP
                            STOP

LL=KK+1

JK=IY(KK)

DENOM(J)=BV(JK)*RATID(KK)

COEFF=(1.-ETA(J))*BV(J)

DO 231 I=1,C

VECTOR(I)=SV(JK,I)*RATIU(KK)

IF(LL.GT.NOFF)GOTU 232

DO 232 KK=LL,NOFF

IF(J.NE.IX(KK))GOTO 232

IF(LORV(KK),EQ.O)GOTO 232

JK=IY(KK)
230
231
                             IF(LORV(KK).EQ.U)GUTU 232

JK=IY(KK)

DENOM(J)=DENOM(J)+BV(JK)*RATIO(KK)

DO 233 I=1,C

VECTOR(I)=VECTOR(I)+SV(JK,I)*RATIO(KK)

CONTINUE

DO 235 KK=1,NOV

IF(J:EQ.JNOSV(KK))GOTO 238

CONTINUE

OFNOM(I)=DENOM(J)+BV(J+1)*RV(J+1)
 233
                              DENOM(J)=DENOM(J)+BV(J+1)*RV(J+1)
DO 237 I=1,C
F(J,I+C)=F(J,I+C)+COEFF*(VECTOR(I)+SV(J+1,I)*RV(J+1))/DENOM(J)
GOTO 240
 235
 237
```

```
238
239
                                      DO 239 I=1,C
F(J,I+C)=F(J,I+C)+CUEFF*VECTOR(I)/DENOM(J)
CONTINUE
240
 : SPECIFICATIONS PART
                                   KK=1
J=JSPC(KK)
JP1=J+1
JM1=J-1
TJ=T(J)
SD0==(1.+SSL(J))
SD1==(1.+SSV(J))
IF(LBL(KK).LT.5)GOTO 260
DO-251 MM=1,NCOND
IF(J.EQ.JCOND(MM))GOTO 255
CONTINUE
DO 252 MM=1,NREBL
IF(J.EQ.JREBL(MM))GOTO 253
CONTINUE
GOTO 260
QQ(J)=0.
CALL ENL(T(J-1),HL)
CALL ENL(TJ,HV)
DO 254 I=1 C
QQ(J)=QQ(J)*SD1
GOTO 260
QQ(J)=QQ(J)*SD1
GOTO 260
CALL ENL(TJ,HL)
IF(ITYPEC(MM).EQ.0)GOTO 256
DO-258 I=1; C
QQ(J)=QQ(J)+(HL(I)-HV(I))*SV(JP1,I)
QQ(J)=QQ(J)+(HL(I)-HV(I))*SV(JP1,I)
GOTO 260
DO 257 I=1; C
QQ(J)=QQ(J)+(HV(I)-HL(I))*SL(J,I)
GQ(J)=QQ(J)*SD0
GOTO(10,20,30,40,50,55,00,65,70,75,80,85,90)LBL(KK)
NDENSER SPECIFICATIONS
                                      KK=1
J=JSPC(KK)
270
251
252
253
254
255
258
 260
 NO CONDENSER SPECIFICATIONS

10 DO 11 MM=1,NCOND

1F(J.EQ.JCOND(MM))GOTO 12

11 CONTINUE

TYPE 906,J

906 FORMAT(1X, --- ERRUR--- '/1X, 'STAGE ', 12, ' IS NOT A CONDENSER')
                                      STOP
QQ(J)=0.
F(J,CT)=0.
CALL ENL(TJ,HL)
CALL ENV(T(J+1),HV)
IF(ITYPEC(MM).EQ.U)GOTO 14
DO-13 I=1,C
QQ(J)=QQ(J)+SV(JP1,I)*(HL(I)-HV(I))
F(J,CT)=F(J,CT)+HL(I)*(KV(JP1)*SV(JP1,I)+SDO*SL(J,I)
1 +SD1*SV(J,I)
QQ(J)=QQ(J)*RV(JP1)
GOTO 265
DO 15 I=1,C
QQ(J)=QQ(J)+(HV(I)-HL(I))*SL(J,I)
F(J,CT)=F(J,CT)+HV(I)*(KV(JP1)*SV(JP1,I)+SDO*SL(J,I))
QQ(J)=QQ(J)*SDO
CALL ENV(TJ,HV)
SUM=0.
DO 16 I=1,C
SUM=SUM+HV(I)*SV(J,I)
F(J,CT)=F(J,CT)+SD1*SUM
                                         STOP
 12
  13
   14
   16
```

```
GOTO 265
   NO REBOILER SPECIFICATIONS

O DO 21 MM=1, NREBL

IF(J.EQ.JREBL(MM))GOTO 22

CONTINUE
IYPE 90/, J

FORMAT(IX, '---ERRUR---'/1X, 'STAGE ', I2, ' IS NOT A REBOILER')
21
907
                           TORMAT(IX, "---ERRUR----/1X, STAGE ', I2, ' IS NOT A REBOILS
STOP

GQ(J)=0.

F(J,CT)=0.

CALL ENL(T(J-1), HL)

CALL ENV(TJ, HV)

IF(ITYPER(MM).EQ.U)GDTD 23

DO 26 I=1;C

QQ(J)=QQ(J)+(HL(I)-HV(I))*SV(J,I)

F(J,CT)=F(J,CT)+HL(I)*(KL(JMI)*SL(JM1,I)+SDO*SL(J,I)+SD1

1 *SV(J,I))

QQ(J)=SD1*QQ(J)

GOTO 265

DO 24 I=1,C

QQ(J)=QQ(J)+(HL(I)-HV(I))*SV(J,I)

F(J,CT)=F(J,CT)+HL(I)*(KL(JMI)*SL(JM1,I)+SD1*SV(J,I))

QQ(J)=SD1*QQ(J)

CALL ENL(TJ,HL)

SUM=0.

DO 25 I=1,C

SUM=SUM+HL(I)*SL(J,I)

F(J,CT)=F(J,CT)+SDO*SUM

GOTO 265
22
26
23
24
25
    CONDENSER-HEAT-DUTY SPECIFIED

O DO 31 MM=1,NCOND

IF (J.EQ.JCOND(MM))GUTO 32

CONTINUE
TYPE 908,J

O8 FORMAT(1X,'---ERRUR---'/1X,'STAGE ',12,' IS NOT A CONDENSER')
31
908
                            F(J,CT)=SPCVAL(KK)
QQ(J)=SPCVAL(KK)
CALL ENV(T(J+1),HV)
CALL ENL(TJ,HL)
 32
                            CALL ENL(TJ, HL)
SUMHV=0.
DO 33 I=1,C
SUMHV=SUMHV+HV(I)*SV(JP1,I)
F(J,CT)=F(J,CT)+RV(JP1)*SUMHV
CALL ENV(TJ, HV)
IF(ITYPEC(MM).EQ.U)GOTO 35
DO 34 I=1;C
F(J,CT)=F(J,CT)+HL(I)*(SD0*SL(J,I)+SD1*SV(J,I))
GOTO 265
SUMLED
33
 34
                              SUML=0.
 35
                             SUMV=0.

DO 36 I=1,C

SUMV=SUMV+HV(I)*SV(J,I)

SUML=SUML+HL(I)*SL(J,I)

F(J,CT)=F(J,CT)+SDO*SUML+SD1*SUMV

GOTO 265
 36
       REBOILER-HEAT-DUTY SPECIFIED

DO 41 MM=1,NREBL

IF(J.EQ.JREBL(MM))GOTO 42

CONTINUE

TYPE 909,J

FORMAT(1X,'---ERROR---'/1X,'STAGE ',I2,' IS NOT A REBOILER')
  40
  41
  909
                               STOP
                              F(J,CT)=SPCVAL(KK)
QQ(J)=SPCVAL(KK)
  42
```

```
CALL ENG(T(J-1), HL)
CALL ENV(TJ, HV)
            DO 43 I=1,C

SUM=SUM+HV(I)*SV(J,I)

F(J,CT)=F(J,CT)+SDI*SUM

IF(ITYPER(MM).EQ.U)GDTO 45

DO 44 I=1;C

F(J,CT)=F(J,CT)+HL(I)*(KL(JM1)*SL(JM1,I)+SDO*SL(J,I))

GOTO 265

SUM=0.
            SUM=0.
            SUM=0.
45
            SUM=0.

SUM1=0.

CALL ENL(TJ,HV)

DO 46 I=1,C

SUM=SUM+HV(I)*SL(J,I)

SUM1=SUM1+HL(I)*SL(JM1,I)

F(J,CT)=F(J,CT)+SDO*SUM+RL(JM1)*SUM1

GOTO 265
REFLUX RATIO SPECIFIED

DO 51 MM=1, NCOND

IF(J.EQ.JCOND(MM))GOTO 52

CONTINUE

TYPE 910, J

FORMAT(1X, "---ERROR---"/1X, "STAGE ", 12," IS NOT A CONDENSER")
             STOP
             F(J,CT)=BL(J)-BV(J)*SPCVAL(KK)
GOTO 265
52
  REBUILER RATIO (V/B) SPECIFIED

DO 56 MM=1,NREBL

IF(J.EQ.JREBL(MM))GOTO 57

CONTINUE
TYPE 911,J
FORMAT(1X,'---ERRUR---'/1X,"STAGE ',12," IS NOT A REBUILER')
55
911
             STOP
             F(J,CT)=BV(J)=BL(J)*SPCVAL(KK)
GOTO 265
57
TEMPERATURE OF THE STAGE IS SPECIFIED F(J,CT)=T(J)-SPCVAL(KK)

GOTO 265
TOTAL VAPUR-FLOW V(J) LEAVING STAGE J IS SPECIFIED L OR "D" IN CASE UF A CONDENSER ]
             F(J,CT)=BV(J)-SPCVAL(KK)
GOTO 265
65
  TOTAL LIQUID-FLUW L(J) LEAVING STAGE J IS SPECIFIED I OR "B" IN CASE OF A REBUILER J
70
             F(J,CT)=BL(J)-SPCVAL(KK)
GOTO 265
VAPOR-MOLE-FRACTION Y(j,1) OF COMPONENT I LEAVING STAGE J SPECIFIED
```

```
L
```

```
L OR Xd(1) IN CASE OF CUNDENSER 1
85
                     IT=ICOMP(KK)
F(J,CT)=SV(J,II)-SPCVAL(KK)*BV(J)
GOTO 265
LIQUID-MOLE-FRACTION X(J,1) OF COMPONENT 1 LEAVING STAGE J SPECIFIED L OR Xb(1) IN CASE OF REBUILER J

90 II=ICOMP(KK)
                      F(J,CT)=SL(J,II)-SPCVAL(KK)*BL(J)
                     KK=KK+1
IF(KK.LE.NSPC)GOTU 270
CONTINUE
265
                     DO 400 J=1,N
WRITE(51,*)J,(F(J,I),I=1,CT)
WRITE(51,*)(QQ(J),J=1,N)
WRITE(52,*)(RV(J),J=1,N)
WRITE(52,*)(RL(J),J=1,N)
WRITE(52,*)(DENOM(J),J=1,N)
400
                     IZ1=47
IZ2=57
                     BL0=200.
RL(1)=200./BL(20)
RL(41)=1.-RL(1)
                     J=1
CALL TRIB(J)
CALL TRIC(J)
CALL OFFC(J,20,T(20))
CALL INVPRT(B)
CALL BCMUL(J)
CALL BFMUL(J)
CALL BAMUL(47,B,COFF)
                     DO 501 JJ=2,9
                      J=JJ
                      JZ=IZ1+JJ-1
CALL TRIA(J)
                      CALL TRIB(J)
                     CALL TRIB(J)
CALL TRIC(J)
CALL APVMOL(J-1)
CALL AQMOL(J-1)
DO 500 1=1;CT
F(J,I)=F(J,I)-VEC1(I)
DO 500 K=1;CP1
B(I,K+C)=B(I,K+C)-ALFA(I,K)
CALL INVPRIB)
CALL BCMUL(J)
CALL BFMUL(J)
500
                       CALL BFMUL(J)
CALL APLMUL(JZ-1)
CALL MBAPL(JZ)
                       CONTINUE
 501
                      J=10
JZ=56
CALL ABSBUT
CALL TRIA(J)
CALL TRIB(J)
CALL APVMUL(J-1)
CALL APVMUL(J-1)
DO 502 I=1,CT
F(J,I)=F(J,I)-VEC1(I)
DO 502 K=1,CP1
B(I,K+C)=B(I,K+C)-ALFA(I,K)
CALL BFMUL(J)
 502
```

```
CALL APLMUL(JZ-1)
CALL MBAPL(JZ)
                                       J=11
                                       JZ=57
CALL ABSTOP
                                       CALL TRIB(J)
CALL TRIB(J)
CALL TRIC(J)
CALL INVPRT(B)
CALL BCMUL(J)
CALL BFMUL(J)
CALL BFMUL(J)
CALL BAMUL(57,B,CUFF)
                                       DO 504 JJ=12,19
J=JJ
                                        JZ=JZ+1
CALL TRIA(J)
                                       CALL TRIA(J)
CALL TRIB(J)
CALL TRIB(J)
CALL TRIC(J)
CALL APVMUL(J-1)
CALL AQMUL(J-1)
DO 506 I=1;CT
F(J,I)=F(J,I)-VEC1(I)
DO 506 K=1;CP1
B(I,K+C)=B(I,K+C)-ALFA(I,K)
CALL INVPRT(B)
CALL BFMUL(J)
CALL BFMUL(J)
CALL MBAPL(JZ-1)
CALL MBAPL(JZ)
CONTINUE
506
                                     J=20
JZ=66
CALL ABSBOT
CALL TRIA(J)
CALL TRIB(J)
CALL AQMUL(J-1)
CALL AQMUL(J-1)
DO 507 I=1;CT
F(J,I)=F(J,I)-VEC1(I)
DO 507 K=1,CP1
B(I,K+C)=B(I,K+C)-ALFA(I,K)
DO 511 I=1;CT
BETA1(I)=Q(10,I)
DO 511 K=1;CP1
BETA2(I,K)=P(56,I,K)
DO 508 KKK=9,1,-1
K=KKK
CALL PVPL(K)
DO 509 I=1;CT
DO 509 L=1;CP1
BETA2(I,L)=P(IZ1+K-1,I,L)-ALFA(I,L)
CALL PVQMUL
CALL PVQMUL
CALL CONTINUE
CALL OFFC(J,1,T(1))
                                        CONTINUE
504
 507
 511
  509
  510
  508
                                          CONTINUE
CALL OFFC(J,1,T(1))
CALL CPLMUL(J,COFF,BETAZ,ALFA)
CALL CQMUL(J)
DO 512 I=1;CT
F(J,I)=F(J,I)-VEC1(I)
DO 512 K=1,CP1
B(I,K+C)=B(I,K+C)-ALFA(I,K)
CALL INVPRT(B)
CALL APLMUL(JZ-1)
CALL BFMUL(JZ-1)
   512
                                            CALL BFMUL(J)
```

```
CALL AQMUL(J-1)
DO 542 I=1;CT
F(J,I)=F(J,I)-VEC1(I)
DO 542 K=1;CP1
B(I,K+C)=B(I,K+C)-ALFA(I,K)
CALL INVPRT(B)
CALL BFMUL(J)
CALL APLMUL(JZ-1)
CALL MBAPL(JZ)
542
                              J=36
CALL
                              CALL CONBC(J,O)
CALL INVPRT(B)
CALL BCMUL(J)
                              CALL BEMUL(J)
                              DO 543 JJ=37,39
                              J=JJ
CALL
                             CALL TRIA(J)
CALL TRIB(J)
CALL TRIB(J)
CALL TRIC(J)
CALL APVMUL(J-1)
CALL AQMUL(J-1)
DD 544 I=1,CT
F(J,I)=F(J,I)-VEC1(I)
DO 544 K=1,CP1
B(I,K+C)=B(I,K+C)-ALFA(I,K)
CALL BFMUL(J)
CALL BFMUL(J)
CONTINUE
                                                 TRIA(J)
544
543
                             J=40

JZ=JZ+1

CALL TRIA(J)

CALL TRIB(J)

CALL TRIC(J)

CALL APYMUL(J-1)

CALL AQMUL(J-1)

DO 545 L=1;CT

F(J,1)=F(J,1)-VEC1(I)

DO 545 K=1,CP1

B(I,K+C)=B(I,K+C)-ALFA(I,K)

CALL INVPRT(B)

CALL BCMUL(J)

DO 546 L=1;CT

BETA1(I)=Q(35,I)

DO 547 KKK=34,21,-1

K=KKK
                              J=40
 545
 546
                               K=KKK
CALL PVUMUL
DO 548 1=1,CT
BETA1(I)=U(K,I)=VEC1(I)
 548
                               CONTINUE

DO 549 I=1,CT

DO 549 K=1,CP1

BETA2(I,K)=P(79,I,K)

DO 550 KKK=34,23,-1
 547
  549
                                K=KKK
KZ=79-K
                                CALL PVPL(K)
DO 551 I=1,CT
DO 551 L=1,CP1
BETAZ(I,L)=P(KZ,I,L)-ALPA(I,L)
  551
550
                                 CONTINUE
                                CALL PVPV(21,22)
CALL OFFC(J,21,T(21))
CALL CPLMUL(J,COFF,BETA2,ALFA)
CALL CQMUL(J)
DO 552 I=1,CT
```

```
F(J,I)=F(J,I)-VEC1(I)
CALL MBAPL(JZ)
CALL BCMUL(J)
CALL BFMUL(J)
552
                                                               J = 41
                                                              JZ≡81
CALL
CALL
                                                            CALL TRIA(J)
CALL TRIB(J)
CALL TRIB(J)
CALL TRIC(J)
CALL OFFA(J,20,T(20))
CALL APVMUL(J-1)
CALL AQMUL(J-1)
CALL AQMUL(J-1)
DO 553 I=1,CT
F(J,I)=F(J,I)-VEC1(I)
DO 553 K=1,CP1
B(I,K+C)=B(I,K+C)-ALFA(I,K)
CALL INVPRT(B)
CALL BCMUL(J)
CALL BCMUL(J-1)
DO 554 I=1;CT
F(J,I)=F(J,I)-VEC1(I)
CALL BFMUL(J)
multiply CC=A.P(80)
multiply CC=A.P(80)
multiply ALFA = AUFF.P(66)
DO 555 K=1,CP1
ALFA(I,K)=CC(I,K+C)+ALFA(I,K)
CALL MBAPL(JZ)
                                                                                                     TRIA(J)
553
554
 555
                                                            JZ=81
DO 556 JJ=42,45
J=JJ
JZ=JZ+1
CALL TRIA(J)
CALL TRIB(J)
CALL TRIC(J)
CALL APVMUL(J-1)
CALL AQMUL(J-1)
DO 557 I=1;CT
F(J,I)=F(J,I)-VEC1(I)
DO 557 K=1,CP1
B(I,K+C)=B(I,K+C)-ALFA(I,K)
CALL INVPRT(B)
CALL BCMUL(J)
CALL BFMUL(J)
CALL BFMUL(J)
CALL APLMUL(JZ-1)
CALL MBAPL(JZ)
CONTINUE
 557
  556
                                                              J=46
JZ=85
CALL TRIA(J)
CALL TRIB(J)
CALL TRIC(J)
CALL APVMUL(J-1)
CALL AQMUL(J-1)
DD 558 I=1;CT
F(J,I)=F(J,I)-VEC1(I)
DD 558 K=1,CP1
B(I,K+C)=B(I,K+C)-ALFA(1,K)
CALL INVPRT(B)
CALL INVPRT(B)
CALL BFMUL(J)
CALL BFMUL(J)
CALL APLMUE(JZ-1)
SUDTTACTION CC=CC-ALFA
CALL MTMUL2(B,CC)
multiply P(46)=B.CC
   558
```

```
J=47
                                                        J=47
CALL REBAB(J,0)
CALL APVMUL(J-1)
CALL AQMUL(J-1)
DO 564 I=1;CT
F(J,I)=F(J,I)-VEC1(I)
DO 564 K=1;CP1
B(I,K+C)=B(I,K+C)-ALFA(I,K)
CALL BFMUL(J)
564
       BACK-SUBSTITUTION
                                                           J=46
                                                           CALL GMPVX(J)
                                                        CALL QMPVX(J)
JZ=85
DD 570 JJ=45,40,-1
J=JJ
CALL QMPVX(J)
CALL QMPLX(J,JZ,47)
JZ=JZ-1
CONTINUE
DD 571 JJ=39,36,-1
J=JJ
CALL QMPVX(J)
570
                                                          J=JJ
CALL QMPVX(J)
CONTINUE
JZ=79
CALL QMPLX(J,JZ,47)
DO 572 JJ=34,23,-1
J=JJ
JZ=JZ-1
CALL QMPVX(J)
CALL QMPVX(J)
CALL QMPLX(J,JZ,47)
CONTINUE
JZ=67
DO 573 JJ=22,21,-1
J=JJ
CONTINUE
J=JJ
 571
 572
   573
                                                           CONTINUE
JZ=66
CALL QMPLX(J,JZ,47)
DO 574 JJ=19,11,-1
J=JJ
JZ=JZ-1
CALL QMPVX(J)
CALL QMPLX(J,JZ,47)
CONTINUE
TY1=20
    574
                                                             IY1=20
JZ=56
CALL GMPLX(J,JZ,20)
DO 575 JJ=9,1,-1
J=JJ
                                                               JZ=JZ-1
                                                              CALL QMPVX(J)
CALL QMPLX(J,JZ,20)
CONTINUE
    575
                                                              STUP
                                                              END
END
SUBROUTINE ENV(TT,HV)
DIMENSION HV(2)
HV(1)=1.
HV(2)=1.
RETURN
                                                             END
SUBROUTINE ENL(TT, HL)
DIMENSION HL(2)
HL(1)=1.
HL(2)=1.
```

```
RETURN
END
                  SUBROUTINE FINDK(TT, AK)
                  DIMENSIUN AK(2)
                  AK(1)=1.
AK(2)=1.
RETURN
END
                  SUMY=0.
DO 30 IP=1,C
                  DELPK=0.
                  IF(K.EQ.IP)DELPK=1.
SUMX=SUMX+XK(I,IP)*(DELPK-SL(J,IP)/BL(J))
SUMY=SUMY+YK(I,IP)*(DELPK-SV(J,IP)/BV(J))
B(I+C,K)=EBYL*(SL(J,I)*SUMX/BL(J)+B(I+C,K))
B(I+C,K+C)=(EBYL*SL(J,I)*SUMY+B(I+C,K+C))/BV(J)
CALL DKBYDT(T(J),AK)
 30
 20
                  CALL DREYDT(T(J),AK,
DO 70 I=1,C
B(I+C,CT)=EBYL*SL(J,I)*AK(I)
IF(ETA(J).EQ.1.)RETURN
FAC1=(1.-ETA(J))/(BV(J+1)*BV(J+1))
DO 80 I=1,C
DO 80 K=1,C
 70
                  DELIK=0.

IF(I.EQ.K)DELIK=1.

CC(I+C,K+C)=FAC1*(DELIK*BV(J+1)-SV(J+1,I))

RETURN
  80
  FOR TOTAL CONDENSERS

100 SQV=1./(BV(J)*BV(J))

SQL=1./(BL(J)*BL(J))

DO 105 I=1,C

E1=-SL(J,I)*SQL
```

```
E2=SV(J,I)*SQV
DO 105 K=1,C
IF(I.EQ.K)GOTO 102
B(I+C,K)=E1
B(I+C,K+C)=E2
GOTO 105
B(I+C,K)=(BL(J)-SL(J,I))*SQL
B(I+C,K+C)=(SV(J,I)-BV(J))*SQV
CONTINUE
RETURN
 102
105
                     RETURN
                     END
                     RETURN
SQL=1./(BL(J)*BL(J))
SQV=1./(BV(J)*BV(J))
DO 60 I=1,C
E1=-SL(J,1)*SQL
E2=SV(J,1)*SQV
DO 60 K=1,C
IF(I.EQ.K)GOTO 51
B(I+C,K)=E1
B(I+C,K+C)=E2
GOTO 60
```

```
B(I+C,K)=(BL(J)-SL(J,I))*SQL
B(I+C,K+C)=(SV(J,I)-BV(J))*SQV
CONTINUE
51
60
                                     RETURN
                                     END
SUBROUTINE TRIA(J)

!SUBROUTINE TO COMPUTE THE ELEMENTS OF THE TRIDIAGONAL "A" MATRICES

INTEGER C,CP1,TWOC,CT

COMMON /C,CP1,TWOC,CT

COMMON /XA/A(9,9)/XCONST/ETA(47),SSL(47),SSV(47)

COMMON /XVAR/SL(47,4),BL(47),RL(47),SV(47,4);BV(47),RV(47),T(47)

DIMENSION HL(4),DHL(4)

DO 10 I=1,CT

DO 10 K=1,CT

A(I,K)=0.

CALL ENL(T(J-1),HL)

CALL ENV(T(J-1),DHL)

SUM=0.

DO 12 I=1,C

A(I,I)=RL(J-1)*HL(I)

A(CT,CT)=RL(J-1)*DHL(I)

A(CT,CT)=RL(J-1,I)*DHL(I)

RETURN

END
SUBROUTINE TRIB(J)

SUBROUTINE TO COMPUTE ELEMENTS OF THE TRIDIAGONAL "B" SUBMATRICES

INTEGER C,CP1,TWOC CT

COMMON /Xb/H(9,9)/XCONST/ETA(47),SSL(47),SSV(47)

COMMON /XVAR/SL(47,4),BL(47),RL(47),SV(47,4);BV(47),RV(47),T(47)

DIMENSION AK(4),XK(4,4),YK(4,4);HL(4),HV(4),DHL(4),DHV(4)

S1=-(1.+SSL(J))

S2=-(1.+SSV(J))

D0 10 1=1,C

B(I,I)=S1

B(I,I)=S1

B(I,I)=S1

B(I,I)=S1

B(I,I)=S1

CALL DKBYDX(T(J),AK)

CALL DKBYDX(T(J),XK)

CALL DKBYDX(T(J),XK)

CALL DKBYDX(T(J),YK)

D0 20 I=1,C

E1=-AK(I)*SL(J,I)

E2=SV(J,I)/BV(J)

D0 20 K=1,C

B(I+C,K)=E1

B(I+C,K)=E1

B(I+C,K)=B(I+C,K)+AK(I)*BL(J)

B(I+C,K)=B(I+C,K)+AK(I)*BL(J)

B(I+C,K)=B(I+C,K+C)=1

SUMX=0.

SUMX=0.

SUMX=0.

SUMX=0.

SUMX=0.

SUMY=0.

SUMY=0.
                                     END
                   SUMY=0.
SUMY=0.
SUMY=0.
"K(1)" VALUES ARE COMPOSITION -INDEPENDENT THEN REMOVE THE "!"
GOTO 24
DO 24 IP=1,C
                                      DELPK=0.

IF(K.EQ.IP)DELPK=1.

SUMX=SUMX+XK(I,IP)*(DELPK-SL(J,IP)*BL(J))

SUMY=SUMY+YK(I,IP)*(DELPK-SV(J,IP)*BV(J))
   24
                                      B(I+C,K)=COEFF*(SL(J,I)*SUMX+B(I+C,K))
B(I+C,K+C)=(EBYL*SL(J,I)*SUMY+B(I+C,K+C))/BV(J)
CALL DKBYDT(T(J),AK)
    20
                                       DO 30 I=1,C
B(I+C,CT)=EBYL*SL(J,I)*AK(I)
```

30

```
CALL ENL(T(J), HL)
CALL ENV(T(J), HV)
CALL DHLDT(T(J), DHL)
CALL DHVDT(T(J), DHV)
                                   SUMX=0.
                                   SUMX=0.

SUMY=0.

DO 33 I=1,C

B(CT,I)=S1*HL(I)

B(CT,I+C)=S2*HV(I)

SUMX=SUMX+SL(J,I)*DHL(I)

SUMY=SUMY+SV(J,I)*DHV(I)

B(CT,CT)=S1*SUMX+S2*SUMY

RETURN
33
                                    END
                                   SUBROUTINE TRIC(J)
'INE TTO CUMPUTE THE ELEMENTS OF THE TRIDIAGONAL "C" MATRICES
INTEGER C,CP1,TWOC,CT
COMMON N,C,CP1,TWOC,CT
COMMON /XC/CC(9,9)/XCUNST/ETA(47),SSL(47),SSV(47)
COMMON /XVAR/SL(47,4),BL(47),RL(47),SV(47,4),BV(47),RV(47),T(47)
COMMON /XLINK/DENUM(47),NOFF,IX(6),IY(6),LORV(6),RATIO(6)
DIMENSION VECTOR(4),HV(4),DHV(4)
DO 10 I=1,CT
DO 10 K=1,CT
CC(I,K)=0.
CALL DHVDT(T(J+1),HV)
CALL DHVDT(T(J+1),DHV)
SUM=0.
SUBROUTINE
10
                                   CALL ENV(T(J+1),HV)
CALL DHVDT(T(J+1),DHV)
SUM=0.
DO 11 I=1,C
CC(I,I+C)=RV(J+1)
CC(CT,I+C)=RV(J+1)*HV(I)
SUM=SUM+SV(J+1,I)*DHV(I)
CC(CT,CT)=RV(J+1)*SUM
IF(ABS(1,-ETA(J)).LE.1.E-6)RETURN
LNK=0
DO 12 I=1.C
11
                                    DO 12 I=1,C

VECTOR(I)=RV(J+1)*SV(J+1,I)

DO 20 KK=1,NOFF

IF((J.EW.IX(KK)).AND.(LURV(KK).EQ.1))GOTO 15

GOTO 20
12
                                      LNK=1
                                     LL=IY(KK)
DO 16 I=1,C
VECTOR(I)=VECTOR(I)+RATLO(LL)*SV(LL,I)
 16
                                    CONTINUE
IF(LNK, EQ.1)GOTU 30
COEFF=(1.~ETA(J))/(BV(J+1)*BV(J+1))
DO 21 I=1,C
E0==COEFF*(BV(J+1,1)
E1=COEFF*(BV(J+1)-SV(J+1,I))
DO 21 K=1;C
IF(I.EQ.K)GUTO 22
CC(I+C,K+C)=E0
GOTO 21
CC(I+C,K+C)=E1
CONTINUE
RETURN
COEFF=(1.~ETA(J))*RV(J+1)/(DENOM(J)*DENOM(J))
DO 31 I=1,C
E0==COEFF*VECTOR(I)
E1=COEFF*(DENOM(J)*VECTUR(I))
DO 31 K=1;C
                                      CONTINUE
  22
   30
                                      DO 31 K=1,C
IF(1,EQ,K)GOTO 32
CC(1+C,K+C)=E0
GOTO 31
CC(1+C,K+C)=E1
CONTINUE
   32
```

```
RETURN
                       END
SUBROUTINE OFFA(J,JEXIT,TT)
SUBROUTINE TO COMPUTE THE ELEMENTS OF (HAVING STRUCTURE OF "A"-SUBMATRICES)
*** Only for Liquid-Interlinks ***
                                                                                                    OF THE OFF-DIAGONAL "A" MATRICES
                      ly for Liquid-Interlinks ***

: stage no. into which the interlink-stream is entering
: stage no. from which the interlink-stream is leaving
: temperature of the interlink-stream entering stage-J
    (TT may be equal to that of the leaving stage-JEXIT
    or TT may be specified)
INTEGER C.CP1.TWOC.CT
COMMON N.C.CP1.TWOC.CT
COMMON /XADFF/AUFF(9,9)
COMMON /XADFF/AUFF(9,9)
COMMON /XLINK/DENUM(47),NOFF,IX(6),IY(6),LORV(6),RATIO(6)
COMMON /XVAR/SL(47,4),BL(47),RL(47),SV(47,4),BV(47),RV(47),T(47)
DIMENSION HL(4);DHL(4)
DO 10 I=1,CT
     J
     JEXIT
TT
                      DO 10 I=1,CT

DO 10 K=1,CT

AOFF(I,K)=0.

DO 20 KK=1,NOFF

IF((J.EQ.IX(KK)).AND.(JEXIT.EQ.IY(KK)))GOTO 30
10
                       CONTINUE
TYPE 900, JEXIT, J
FORMAT(1X, There is no stream from stage', 12, 'to stage', 12)
20
900
                      STOP

IF (LORV(KK), EQ.0)GOTO 40

TYPE 901, J

FORMAT(1X, subroutine OFFA is strictly for liquid interlinks
1 and should not be called for vapor streams. Check stage ;12)
30
901
                      STOP
CALL ENL(TT, HL)
DO 50 I=1, C
AOFF(I, I)=RATIO(KK)
AOFF(CT, I)=RATIO(KK)*HL(I)
IF(ABS(TT-T(JEXIT)).GT.1,E-6)RETURN
CALL DHLDT(TT, DHL)
DO 70 I=1, C
AOFF(CT, CT)=AOFF(CT, CT)+SL(JEXIT, I)*DHL(I)
AOFF(CT, CT)=AOFF(CT, CT)*RATIO(KK)
RETURN
END
40
50
70
                       END
 SUBRUUTINE OFFC(J,JEXIT,TT)
SUBROUTINE TO COMPUTE THE ELEMENTS OF THE OFF-DIAGONAL "C" MATRICES (HAVING STRUCTURE OF "C"-SUBMATRICES)
                       *** Only
                                          Vapor-Interlinks ***
 *******
      JEXIT
  10
  20
```

900

STOP

```
IF(LORV(KK).EQ.1)GOTO 40
TYPE 901,J
FORMAT(1X, Subroutine OFFC is strictly for vapor interlinks
1 and should not be called for liquid streams. Check stage, 12)
30
901
40
                                 CALL ENV(TT, HV)

DD 50 I=1, C

COFF(I, I+C)=RATIO(KK)

COFF(CT, I+C)=RATIO(KK)*HV(I)

IF(ABS(TT-T(JEXIT)).GT.1.E-6)GOTO 70

CALL DHVDT(TT, DHV)

DD 60 I=1, C

COFF(CT, CT)=COFF(CT, CT)+SV(JEXIT, I)*DHV(I)

COFF(CT, CT)=RATIO(KK)*CUFF(CT, CT)

IF(ABS(1.-ETA(J)).LE.1.E-6)RETURN

DO 80 I=1, C

VECTOR(I)=RV(J+1)*SV(J+1, I)

DO 90 LL=1, NOFF

IF((J.EQ.IX(LL)).AND.(LURV(LL).EQ.1))GOTO 95

GOTO 90

DO 99 I=1, C
                                   CALL ENV(TT, HV)
50
60
70
80
                                  GOTO 90
DO 99 I=1,C
VECTOR(I)=VECTOR(I)+RATIO(LL)*SV(IY(LL),I)
CONTINUE
FAC1=(1.-ETA(J))*RATIO(KK)/DENOM(J)
DO 100 I=1,C
E0=-FAC1*VECTOR(I)/DENOM(J)
E1=FAC1*VECTOR(I)/DENOM(J)
DO 100 K=1,C
IF(I.EQ.K)GOTO 110
COFF(I+C,K+C)=E0
GOTO 100
COFF(I+C,K+C)=E1
CONTINUE
RETURN
95
99
 110
 100
                                    RETURN
                                   END
 SUBROUTINE PROB(J,LABEL,ITYPE, VAL,ICOM)

This subroutine computes the last row of the "A", "B" & "C" submatrices, depending on the type of specification variable(LABEL) for any stage J : stage no.
                                     : stage no.
                                             if stage J is a condenser or a reboiler then set ITYPE = U for a "partial" condenser or reboiler
           ITYPE
                                   then set ITYPE = 0 for a "partial" condenser or reboiler else assign any integer to ITYPE if no specification-value is required then assign any real number to VAL else assign given value of the specification variable to VAL: component no.(if not required then assign any integer) INTEGER C.CP1.TWOC.CT COMMON N.C.CP1.TWOC.CT COMMON /XA/A(9,9)/XB/B(9,9)/XC/CC(9,9) COMMON /XAR/SL(47,4),BL(47),RL(47),SV(47,4),BV(47),RV(47),T(47) COMMON /XCONST/ETA(47),SSL(47),SSV(47),DHV(4),DHL(4) COMMON /XCONST/ETA(47),SSL(47),SSV(47),DHV(4),DHL(4) GOTO(10,20,30,40,50,60,70,80,90,100,110,120,130)LABEL TYPE 900,J,LABEL FORMAT(1X, error in specification, for stage', I3, 'spec. type', I3) STOP
          VAL
          ICOM
   900
                                      STOP
          NO CONDENSER SPECIFICATIONS
                                     S1=-(1.+SSL(J))
S2=-(1.+SSV(J))
IF(ITYPE.EQ.1)GOTU 15
CALL ENV(T(J),HV)
CALL DHVDT(T(J),DHV)
   10
                                     SUM=0.

DO 12 I=1,C

B(CT,I)=S1*HVP1(I)

B(CT,I+C)=S2*HV(I)
```

```
SUM=SUM+SV(J,I)*DHV(I)
B(CT,CT)=S2*SUM
CALL DHVDT(T(J+1),DHV)
CC(CT,CT)=0.
DO 13 I=1,C
CC(CT,I)=0.
CC(CT,I)=0.
CC(CT,I+C)=RV(J+1)*HVP1(I)
CC(CT,CT)=CC(CT,CT)+(RV(J+1)*SV(J+1,I)+S1*SL(J,I))*DHV(I)
RETURN
CALL FNL(T(J) HL)
12
13
                                      RETURN

CALL ENL(T(J), HL)

CALL DHLDT(T(J), DHL)

B(CT,CT)=0.

DO 16 I=1, C

B(CT,I)=S1*HL(I)

B(CT,I+C)=S2*HL(I)

B(CT,CT)=B(CT,CT)+(RV(J+1)*SV(J+1,I)+S2*SV(J,I)+S1*SL(J,I))*

1 DHL(I)

CC(CT,I)=0.

CC(CT,I+C)=RV(J+1)*RV(J+1)*HL(I)

CC(CT,CT)=0.

RETURN

OLER SPECIFICATIONS
15
16
               RETURN

REBOILER SPECIFICATIONS

S1=-(1.+SSL(J))

S2=-(1.+SSV(J))

IF(ITYPE.EQ.1)GOTU 25

CALL ENL(T(J-1), HLM1)

CALL ENL(T(J), HL)

CALL DHLDT(T(J-1), DHL)

A(CT,CT)=0.

DO 22 I=1,C

A(CT,I)=RL(J-1)*HLM1(I)

A(CT,I+C)=0.

A(CT,CT)=A(CT,CT)+(RL(J-1)*SL(J-1,I)+S2*SV(J,I))*DHL(I)

B(CT,I)=S1*HL(I)

B(CT,I+C)=S2*HLM1(I)

CALL DHLDT(T(J),DHL)

SUM=0.
1NO
20
                                        SUM=0.

DO 23 I=1,C

SUM=SUM+SL(J,I)*DHL(I)

B(CT,CT)=S1*SUM

RETURN
 23
                                        RETURN

CALL ENL(T(J-1), HLM1)

CALL DHLDT(T(J-1), DHL)

A(CT,CT)=0.

DO 26 I=1, C

A(CT,I)=RL(J-1)*HLM1(I)

A(CT,I)=0.

A(CT,CT)=A(CT,CT)+(RL(J-1)*SL(J-1,I)+S1*SL(J,I)+S2*SV(J,I))*

1 DHL(I)

B(CT,I)=S1*HLM1(I)

B(CT,I)=S1*HLM1(I)

B(CT,I)=S2*HLM1(I)

B(CT,CT)=0.

RETURN

VSER-HEAT-DUTY SPECIFIED
 25
  26
          RÈTURN

CONDENSER-HEAT-DUTY SPECIFIED

$1=-(1.+$SL(J))

$2=-(1.+$SV(J))

CALL ENV(T(J+1),HV)

CALL DHVDT(T(J+1),DHV)

SUM=0.

DO 31 I=1,C

CC(CT,I)=0.

CC(CT,I+C)=RV(J+1)*HV(I)

SUM=SUM+SV(J+1,I)*DHV(I)

CC(CT,CT)=RV(J+1)*SUM

IF(ITYPE.EQ.1)GOTU 35

CALL ENL(T(J),HL)

CALL DHLDT(T(J),DHL)
   30
    31
```

```
CALL DHVDT(T(J),DHV)
E1=0.
E2=0.
DO 32 I=1,C
B(CT,I)=S1*HL(I)
B(CT,I+C)=S2*HV(I)
E1=E1+SL(J,I)*DHL(I)
E2=E2+SV(J,I)*DHV(I)
B(CT,CT)=S1*E1+S2*E2
RETURN
CALL ENL(T(J).HL)
32
                                                         CALL ENL(T(J), HL)
CALL DHLDT(T(J), DHL)
35
                                                       CALL DHLDT(T(J),DHL)
E1=0.
E2=0.
DD 36 I=1,C
B(CT,I)=S1*HL(I)
B(CT,I+C)=S2*HL(I)
E1=E1+SL(J,I)*DHL(I)
E2=E2+SV(J,I)*DHL(I)
B(CT,CT)=S1*E1+S2*E2
RETURN
EP-HEAT-DUTY SPECIFI
36
           REBOILER-HEAT-DUTY SPECIFIED

S1=-(1.+SSL(J))

S2=-(1.+SSV(J))

CALL ENL(T(J-1), HLM1)

CALL DHLDT(T(J-1), DHL)
 40
                                                      CALL ENL(T(J=1), HLM1)
CALL DHLDT(T(J=1), DHL)
SUM=0.
DO 41 I=1, C
A(CT, I)=RL(J=1)*HLM1(I)
A(CT, I+C)=0.
SUM=SUM+SL(J=1, I)*DHL(I)
A(CT, CT)=RE(J=1)*SUM
A(CT, CT)=RE(J=1)*SUM
A(CT, CT)=RE(J=1)*SUM
CALL ENL(T(J), HL)
CALL ENL(T(J), HL)
CALL ENL(T(J), DHL)
CALL DHLDT(T(J), DHL)
CALL DHLDT(T(J), DHL)
CALL DHLDT(T(J), DHL)
E1=0.
E2=0.
B(CT, I)=S1*HL(I)
B(CT, I+C)=S2*HV(I)
E1=E1+SL(J, I)*DHL(I)
E2=E2+SV(J, I)*DHL(I)
E1=E1+SL(J, I)*DHL(I)
B(CT, CT)=S1*E1+S2*E2
RETURN
CALL ENV(T(J), HV)
CALL DHVDT(T(J), DHV)
SUM=0.
DO 46 I=1, C
B(CT, I)=S1*HLM1(I)
B(CT, I+C)=S2*SUM
RETURN
X RATIO (L/D) SPECIFIED
DO 51 I=1, C
 41
  42
  45
    46
                RETURN

REFLUX RATIO (L/D) SPECIFIED

DO 51 I=1,C

B(CT,I)=1.

B(CT,I+C)=-VAL

B(CT,CT)=0.

DO 52 I=1,CT

CC(CT,I)=0.

RETURN

REBOILER RATIO SPECIFIED
    50
     51
     52
                 REBOILER RATIO SPECIFIED

DO 61 I=1,C

B(CT,I)=-VAL

B(CT,I+C)=1.

B(CT,CT)=0.
      60
      61
```

```
DO 62 I=1,CT
A(CT,I)=0.
RETURN
SPECIFIED
DO 71 I=1,CT
A(CT,I)=0.
B(CT,I)=0.
CC(CT,I)=0.
B(CT,CT)=1.
RETURN
SPECIFIED (CT
62
      T(j)
70
71
                    RETURN

SPECIFIED (or D specified for condensers)

DO 81 I=1;CT
A(CT,I)=0.
CC(CT,I)=0.
DO 82 I=1,C
B(CT,I)=0.
B(CT,I+C)=1.
B(CT,CT)=0.
RETURN

SPECIFIED (or B specified for reboilers)
DO 91 I=1;CT
A(CT,I)=0.
CC(CT,I)=0.
CC(CT,I)=0.
B(CT,I+C)=0.
B(CT,I+C)=0.
B(CT,I+C)=0.
B(CT,I+C)=0.
B(CT,I+C)=0.
RETURN

1,1) SPECIFIED (or d(1) specified for condense
1 V(j)
80
81
82
 1 L(j)
91
92
RETURN

SPECIFIED (or xd(i) specified for condensers)

DO 121 1=1,CT

A(CT,I)=0.

CC(CT,I)=0.

DO-122 I=1,C

B(CT,I)=0.

B(CT,I)=0.

B(CT,I)=0.

B(CT,C+ICOM)=B(CT,C+ICOM)+1.

B(CT,CT)=0.

RETURN

SPECIFIED (or xb(i) specified for reboilers)
 121
 122
                            RETURN
SPECIFIED (or xb(i) specified for reboilers)
DO 131 I=1,CT
A(CT,I)=0.
CC(CT,I)=0.
DO 132 I=1,C
B(CT,I)=-VAL
B(CT,I+C)=0.
B(CT,ICUM)=B(CT,ICUM)+1.
B(CT,CT)=0.
RETURN
FND
      x(j,i)
 130
 131
 132
                             END
                             SUBROUTINE PVPL(JPV)
COMMON N,C,CP1,TWUC,CT
COMMON /XP/P(85,9,5)/XBETA2/BETA2(9,5)/XALFA/ALFA(9,5)
DO 10 I=1,CT
```

```
DO 10 K=1,CP1
ALFA(I,K)=0.0
DO 10 L=1,CP1
ALFA(I,K)=ALFA(I,K)+P(JPV,I,L)*BETA2(L+C,K)
10
                             RETURN
                            END
SUBROUTINE PVPV(JPV1,JPV2)
INTEGER C,CP1,TWOC,CT
COMMON N,C,CP1,TWOC,CT
COMMON /XP/P(85,9,5)/XALFA/ALFA(9,5)
DO 10 I=1,CT
DO 10 K=1,CP1
ALFA(I,K)=0.0
DO 10 L=1,CP1
ALFA(I,K)=ALFA(I,K)+P(JPV1,I,L)*P(JPV2,L+C,K)
RETURN
END
                             END
10
                             END
                            SUBROUTINE PLPV(U1,U2,U3)
INTEGER C,CP1,TWOC,CT
COMMON N,C,CP1,TWUC,CT
DIMENSION U1(9,5),U2(9,5),U3(9,5)
DO 10 I=1,CT
DO 10 K=1,CP1
U3(I,J)=U1(I,CT)*U2(CT,K)
DO 10 L=1,C
U3(I,K)=U3(I,K)+U1(I,L)*U2(L,K)
RETURN
END
10
                             END
                            END

SUBROUTINE PLPL(U1,U2,U3)

INTEGER C,CP1,TWOC,CT

COMMON N,C,CP1,TWOC,CT

DIMENSION U1(9,5),U2(9,5),U3(9,5)

DO 10 I=1,CT

DO 10 K=1,CP1

U3(I,J)=U1(I,CT)*U2(CT,K)

DO 10 L=1,C

U3(I,K)=U3(I,K)+U1(I,L)*U2(L,K)

RETURN

END
 10
                            END
SUBROUTINE APVMUL(J)

INTEGER C,CP1,TWOC,CT

COMMON N,C,CP1,TWUC,CT

COMMON /XA/A(9,9)/XP/P(85,9,5)/XALFA/ALFA(9,5)

RRR=A(1,1)

IF(ABS(RRR-1.0).LE.1.0E-06) GU TO 30

DO 10 I=1,C

DO 10 K=1,CP1

ALFA(I,K)=RRR*P(J,I,K)

GD TO 40

DO 20 K=1,CP1

ALFA(I,K)=P(J,I,K)

DO 50 K=1,CP1

ALFA(I,K)=P(J,I,K)

DO 50 K=1,CP1

ALFA(CT,K)=A(CT,CT)*P(J,CT,K)

DO 50 L=1,C

ALFA(CT,K)=ALFA(CT,K)+A(CT,L)*P(J,L,K)

RETURN
                              END
 10
  30
 20
  40
  50
                              RETURN
                              END
SUBROUTINE APLMUL(JZM1)
INTEGER C,CP1,TWOC,CT
COMMON N,C,CP1,TWOC,CT
COMMON /XA/A(9,9)/XP/P(85,9,5)/XALFA/ALFA(9,5)
RRR=A(1,1)
IF(ABS(1.0-RRR).LE.1.E-U6) GD TO 20
DO 10 I=1.C
DO 10 K=1,CP1
```

```
ALFA(I,K)=RRR*P(JZM1,I,K)
GO TO 50
          10
                                       DO 30 K=1,CP1
ALFA(I,K)=P(JZM1,I,K)
DO 60 K=1,CP1
ALFA(CT,K)=A(CT,CT)*P(JZM1,CT,K)
DO 60 L=1,C
ALFA(CT,K)=ALFA(CT,K)+A(CT,L)*P(JZ,L,K)
RETURN
FND
          20
          30
          50
          60
                                        RETURN
END
SUBROUTINE CPVMUL(J, COFF, BETA2, ALFA)
INTEGER C, CP1, TWOC, CT
COMMON N, C, CP1, TWOC, CT
DIMENSION COFF(9,9), BETA2(9,5), ALFA(9,5)
RRR=COFF(1, CP1)
IF(ABS(1.U-RRR).LE.1.UE-06) GO TO 30
DO 10 I=1, C
DO 10 K=1, CP1
ALFA(I, K)=RRR*BETA2(I+C, K)
GO TO 40
          10
                                       DO 20 I=1,C

DO 20 K=1,CP1

ALFA(I,K)=BETA2(I+C,K)

IF(ABS(1.0-ETA(J)).LE.1.0E-06) GO TO 70

DO 50 I=CP1

ALFA(I,K)=0.0

DO 50 L=CP1,TWOC

ALFA(I,K)=ALFA(I,K)+COFF(I,L)*BETA2(L,K)

GO TO 80

DO 60 I=CP1,TWOC
           30
          20
50
                                       GO TO 80
DO 60 I=CP1, TWDC
DO 60 K=1, CP1
ALFA(I,K)=0.0
DD 90 K=1, CP1
ALFA(CT,K)=0.0
DO 90 L=CP1, CT
ALFA(CT,K)=ALFA(CT,K)+CUFF(CT,L)*BETA2(L,K)
RETURN
           70
           60
 80
                                       ALFA(CT,K)=ALFA(CT,K)+CUFF(CT,L)*BETA2(L,
RETURN
END
SUBROUTINE CPLMUL(J,COFF,BETA2,ALFA)
INTEGER C,CP1,TWOC,CT
COMMON N,C,CP1,TWOC,CT
DIMENSION COFF(9,9),BETA2(9,5),ALFA(9,5)
RRR=COFF(1,CP1)
IF(ABS(1,U-RRR),LE.1.OE-O6) GO TO 30
DO 10 I=1,C
DO 10 K=1,CP1
ALFA(I,K)=RRR*BETA2(I+C,K)
GO TO 40
DO 20 I=1,C
DO 20 K=1,CP1
ALFA(I,K)=BETA2(I+C,K)
IF(ABS(1,U-ETA(J)),LE.1.E-O6) GO TO 70
DO 50 I=CP1,TWOC
DO 50 L=CP1,TWOC
ALFA(I,K)=0.0
DO 50 L=CP1,TWOC
ALFA(I,K)=ALFA(I,K)+COFF(I,L)*BETA2(L,K)
GO TO 80
DO 60 I=CP1,TWOC
           90
            10
            30
            20
            40
             50
                                          DO 60 I=CP1,TWOC
DO 60 K=1,CP1
ALFA(I,K)=0.0
DO 90 K=1,CP1
ALFA(CT,K)=0.0
DO 90 L=CP1,TWOC
             70
             60
             80
```

```
ALFA(CT,K)=ALFA(CT,K)+CDFF(CT,L)*BETA2(L,K)
RETURN
   90
                         END
                        SUBROUTINE BCMUL(J)
INTEGER C.CP1, TWOC, CT
COMMON N.C.CP1, TWOC, CT
COMMON /XB/B(9,9)/XCC/CC(9,9)/XP/P(85,9,5)
                        COMMON /XB/B(9,9)/XCC/CC(9,9)/XP/P(85,9)
RRR=CC(1,C+1)
CTCT=CC(CT,CT)
IF(ABS(1.0-ETA(J)).LE.1.E-06) GO TO 100
IF(ABS(1.0-RRR).LE.1.E-06)GO TO 110
DO 10 I=1,CT
P(J,I,CP1)=B(I,CT)*CTCT
DO 10 K=1,C
P(J,I,K)=RRR*B(I,K)
DO 10 L=CP1,CT
P(J,I,K)=P(J,I,K)+B(I,L)*CC(L,K+C)
RETURN
DO 20 I=1.CT
   10
                        RETURN
DO 20 I=1,CT
P(J,I,CP1)=B(I,CT)*CTCT
DO 20 K=1,C
P(J,I,K)=B(I,K)
DO 20 L=CP1,CT
P(J,I,K)=P(J,I,K)+B(I,L)*CC(L,K+C)
RETURN
IF(ABS(1,G=PPP), LE 1,E=U6), CD TO 1
   110
   20
                        IF(ABS(1.0=RRR).LE.1.E=06) GD TO 120
DO 30 I=1,CT
P(J,1,CP1)=B(I,CT)*CTCT
DO 30 K=1,C
P(J,I,K)=RRR*B(I,K)
RETURN
   100
30
                        DO 40 I=1,CT
P(J,I,CT)=B(I,CT)*CTCT
DO 40 K=1,C
P(J,I,K)=B(I,K)
RETURN
   120
   40
                         END
                         SUBROUTINE BAMUL(JZ, B, COFF)
INTEGER C, CP1, TWOC, CT
COMMON N, C, CP1, TWUC, CT
COMMON /XP/P(85,9,5)
                         DIMENSION COFF(9,9),B(9,9)

RRR=COFF(1,1)

CTCT=COFF(CT,CT)

IF(ABS(1.0-RRR).LE.1.0E-06) GO TO 100

DO 10 I=1,CT

P(JZ,I,CP1)=B(I,CT)*CTCT

DO 10 K=1,C

P(JZ,I,K)=RRR*B(I,K)+B(I,CT)*COFF(CT,K)

RETURN
    10
                         DO 20 I=1,CT
P(JZ,I,CP1)=B(I,CT)*CTCT
DO 20 K=1,C
P(JZ,I,K)=B(I,K)+B(I,CT)*COFF(CT,K)
RETURN
    100
    20
                         END
SUBROUTINE MBAPL(JZ)
INTEGER C,CP1,TWOC,CT
COMMON /XP/B(9,9)/XALFA/ALFA(9,5)/XP/P(85,9,5)
DO 10 I=1,CT
DO 10 K=1,CP1
P(JZ,I,K)=-B(I,CT)*ALFA(CT,K)
DO 10 L=1,C
P(JZ,I,K)=P(JZ,I,K)-B(I,L)*ALFA(L,K)
RETURN
END
      10
```

END

```
SUBROUTINE MBAPV(JZ)
INTEGER C,CP1,TWOC,CT
COMMON N,C,CP1,TWOC,CT
COMMON /XB/B(9,9)/XALFA/ALFA(9,5)/XP/P(85,9,5)
                                                            DO 10 I=1,CT

DO 10 K=1,CP1

P(JZ,I,K)=-B(I,CT)*ALFA(CT,K)

DO 10 L=1,C

P(JZ,I,K)=P(JZ,I,K)-B(I,L)*ALFA(L,K)

RETURN
       10
                                                             END
                                                            SUBROUTINE AQMUL(JM1)
INTEGER C,CP1,TWOC,CT
COMMON N,C,CP1,TWOC,CT
COMMON /XA/A(9,9)/XO/Q(47,9)/XVEC1/VEC1(9)
                                                             RRR=A(1,1)
VEC1(CT)=A(CT,CT)*Q(JM1,CT)
                                                         VEC1(CT)=A(CT,CT)*Q(JM1,CT)
SUM=0.0
IF(ABS(1.0-RRR).LE.1.E-U6) GD TO 100
DO 10 I=1,C
VEC1(I)=RRR*Q(JM1,I)
VEC1(I+C)=0.0
SUM=SUM +A(CT,I)*Q(JM1,I)
VEC1(CT)=VEC1(CT)+SUM
RETURN
DO 20 I=1,C
VEC1(I)=Q(JM1,I)
VEC1(I+C)=0.0
SUM=SUM+A(CT,I)*Q(JM1,I)
VEC1(CT)=VEC1(CT)+SUM
RETURN
PO 20 I=1,C
VEC1(I)=Q(JM1,I)
VEC1(I)=Q(JM1,I)
VEC1(I)=Q(JM1,I)
VEC1(CT)=VECI(CT)+SUM
RETURN
END
       10
       100
       20
                                                        END
SUBROUTINE COMUL(J)
INTEGER C,CP1,TWOC,CT
COMMON N,C,CP1,TWOC,CT
COMMON N,C,CP1,TWOC
VEC1(CT)=COFF(CT,CT)*BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1/BETA1
                                                             END
                                                           VEC1(I+C)=VEC1(I+C)+COFF(I+C,L)*BETA1(L)
RETURN
DO 20 I=1,C
VEC1(I)=BETA1(I+C)
VEC1(I+C)=0.0
VEC1(CT)=VEC1(CT)+COFF(CT,I+C)*BETA1(I+C)
DO 20 L=CP1,TWOC
VEC1(I+C)=VEC1(I+C)+COFF(I+C,L)*BETA1(L)
RETURN
IF(ABS(1.0-RRR).LE.1.E-U6) GD TO 150
DO 30 I=1,C
VEC1(I)=RRR*BETA1(I+C)
VEC1(I+C)=0.0
VEC1(CT)=VEC1(CT)+COFF(CT,I+C)*BETA1(I+C)
RETURN
10
         50
                 20
 100
           30
                                                              RETURN
DO 40 I=1,C
VEC1(I)=BETA1(I+C)
VEC1(I+C)=0.0
VEC1(CT)=VEC1(CT)+COFF(CT,I+C)*BETA1(I+C)
RETURN
END
SUBBOURTINE
           150
            40
                                                                 SUBROUTINE BEMUL(J)
INTEGER C, CP1, TWOC, CT
```

```
COMMON N.C.CP1.TWUC.CT
COMMON /XB/B(9,9)/XF/F(85,9)/XQ/Q(85,9)
                        DO 10 I=1,CT
Q(J,I)=0.0
DO 10 K=1,CT
                        Q(J,I)=B(I,K)*F(J,K)
RETURN
    10
                         END
                        END
SUBROUTINE QMPVX(J)
INTEGER C.CP1.TWOC.CT
COMMON N.C.CP1.TWOC.CT
COMMON /XP/P(85,9,5)/XQ/Q(47,9)
DO 10 I=1.CT
DO 10 K=1,CP1
                        Q(J,I)=Q(J,I)-P(J,I,K)*Q(J+1,K+C)
RETURN
  10
                         END
                        SUBROUTINE QMPLX(J,JP,JQ)
INTEGER C,CP1,TWOC,CT
COMMON N,C,CP1,TWOC,CT
COMMON /XP/P(85,9,5)/XQ/Q(47,9)
                        DO 10 I=1,CT
Q(J,I)=Q(J,I)-P(JP,I,CP1)*Q(JQ,CT)
DO 10 K=1,C
Q(J,I)=Q(J,I)-P(JP,I,K)*Q(JQ,K)
RETURN
     10
                       SUBROUTINE INVPRT(B)

INTEGER C,CT

COMMON C,CT,N

DIMENSIUN B(CT,CT),BS(4,4),BS1(4,4),PA(4),PB(4)

DO 1 I=1,C

PA(I)=1./B(I,I)

CONTINUE

DO 2 J=1,C

BS(I,J)=PA(I)*B(I,J+C)

CONTINUE

DO 3 J=1,C

DO 3 J=1,C

DO 3 J=1,C

BS1(I,J)=BS1(I,J)+B(I+C,IJ)*BS(IJ,J)

CONTINUE

DO 4 I=1,C
                         END
                         CONTINUE
DO 4 I=1,C
DO 4 J=1,C
BS1(I,J)=B(I+C,J+C)=BS1(I,J)
CONTINUE
CALL MATIN(BS1,C,XXX,0,DETERM)
DO 5 I=1,C
DO 5 J=1,C
B(I+C,J+C)=BS1(I,J)
BS1(I,J)=B(I+C,J)*PA(J)
B(I,J+C)=0.
B(I,J+C)=0.
CONTINUE
DO 6 I=1,C
4
                          CONTINUE

DO 6 I=1,C

DO 6 J=1,C

B(I,J)=0.

DO 6 IJ=1,C

B(I,J+C)=B(I,J+C)-BS(I,IJ)*B(IJ+C,J+C)

CONTINUE

DO 7 I=1,C

DO 7 J=1,C

BS(I,J)=0.
6
                           DO 7 J=1,C
BS(1,J)=0.
DO 7 IJ=1,C
B(I+C,J)=B(I+C,J)=B(I+C,C+IJ)*BS1(IJ,J)
```

1

2

3

```
B(I,J)=B(1,J)=B(I,IJ+C)*BS1(IJ,J)
CONTINUE
DO 9 I=1,C
BS(I,I)=PA(1)
PA(I)=0.
DO 9 J=1,C
B(I,J)=B(I,J)+BS(1,J)
CONTINUE
DO 10 I=1,C
DO 10 II=1,C
PA(I)=PA(I)+B(I,II)*B(II,CT)+B(I,II+C)*B(II+C,CT)
PB(I)=PB(I)+B(I+C,II)*B(II,CT)+B(I+C,II+C)*B(II+C,CT)
CONTINUE
7
9
10
                                   CONTINUE
                                  DO 11 I=1,C
BB=BB+B(CT,I)*PA(I)+B(CT,I+C)*PB(I)
                                DO 11 I=1,C
   BB=BB+B(CT,I)*PA(I)+B(CT,I+C)*PB(I)

CONTINUE
B(CT,CT)=1./(B(CT,CT)-BB)

DO 12 I=1,C
B(I,CT)=-PA(I)*B(CT,CT)
B(I+C,CT)=-PB(I)*B(CT,CT)
PA(I)=0.

CONTINUE
DO 13 I=1,C
PA(J)=PA(J)+B(CT,II)*B(II,J)+B(CT,II+C)*B(II+C,J)
PB(J)=PB(J)+B(CT,II)*B(II,J+C)+B(CT,II+C)*B(II+C,J+C)

CONTINUE
DO 14 I=1,C
B(CT,I)=-B(CT,CT)*PA(I)
B(CT,I+C)=-B(CT,CT)*PA(I)
B(CT,I+C)=-B(CT,CT)*PB(I)

CONTINUE
DO 15 J=1,C
B(I,J)=B(I,J)-B(I,CT)*PA(J)
B(I,J+C)=B(I,J+C)-B(I,CT)*PB(J)
B(I,J+C)=B(I,J+C)-B(I+C,CT)*PB(J)
B(I+C,J+C)=B(I+C,J)-B(I+C,CT)*PB(J)
B(I+C,J+C)=B(I+C,J+C)-B(I+C,CT)*PB(J)
B(I+C,J+C)=B(I+C,J+C)-B(I+C,CT)*PB(J)
B(I+C,J+C)=B(I+C,J+C)-B(I+C,CT)*PB(J)
B(I+C,J+C)=B(I+C,J+C)-B(I+C,CT)*PB(J)
BS1(I,J)=0

CONTINUE
BB=0
DO 22 I=1,C
11
12
13
14
15
                                  BB=0.

DD 22 I=1,C

PB(I)=0.

CONTINUE
22
                                    RETURN
                                    END
                                   SUBROUTINE MATIN(A,N,B,M,DETERM)
A=CO-EFFICIENT OF ORDER N
B=VECTOR OF ORDER N
M=IF M IS SET TO ZERO, ONLY INVERSEIS COMPUTED
DETERM=VALUE OF DETERMENENT RETURNED
                                    DIMENSION A(N,N), B(N,1), IPIVOT(70), INDEX(70,2), DT(70) EQUIVALENCE (IROW, JROW), (ICOLOM, JCOLUM), (AMAX, T, SWAP)
 C
                                    INITIALIZATION
                                    DETERM=1.0
DO 20 J=1,N
IPIVOT(J)=0
SEARCH FOR PIVOT ELEMENT
DO 550 1=1,N
 20
C
                                    AMAX=0.0

DO 105 J=1,N

IF(IPIVOT(J)-1)60,105,60

DO 100 K=1,N

IF(IPIVOT(K)-1)80,100,740

IF(AMAX-ABS(A(J,K)))85,100,100
 60
 80
 85
                                      IROW=J
                                     ICULUM=K
```

```
AMAX=ABS(A(J,K))
CONTINUE
CONTINUE
IPIVOT(ICOLUM)=IPIVOT(ICOLUM)+1
INTRECHANGE ROWS TO PUT PIVOT VECTORS ONDIAGONAL
IF(IROw-ICOLUM)140,260,140

DETERM=DETERM
DO 200 L=1, N
SWAP=A(IRUM,L)
A(IRUW,L)=A(ICOLUM,L)
A(IRUW,L)=SWAP
IF(M)260,260,210

DO 250 L=1,M
SWAP=B(IRUW,L)
B(IROW,L)=SWAP
INDEX(I,1)=IROW
INDEX(I,1)=IROW
INDEX(I,1)=IROW
INDEX(I,2)=ICOLUM
DIVIDE PIVOT ROW BY PIVOR ELEMENT
PIVOT=A(ICOLUM,ICOLUM)
DT(I)=PIVUT
A(ICOLUM,L)=A(ICOLUM,L)/PIVOT
REDUCE NON PIVOT ROWS
IF(M)380,380,360
DO 370 L=1,M
B(ICOLUM,L)=B(ICOLUM,L)/PIVOT
DO 550 L1=1,N
B(ICOLUM,L)=B(ICOLUM,L)/PIVOT
DO 550 L1=1,N
B(ICOLUM,L)=B(ICOLUM,L)/PIVOT
DO 550 L1=1,N
A(L1,ICOLUM)=0.0
A(L1,ICOLUM)=0.0
A(L1,L)=A(I,L)-A(ICOLUM,L)*T
IF(M)550,550,460
DO 500 L=1,M
B(L1,L)=B(L1,L)-B(ICOLUM,L)*T
CONTINUE
INTERCHANGE THE CULUMNS
DO 710 I=1,N
L=N+1-I
IF(INDEX(L,1)-INDEX(L,2))630,710,630
JROW=INDEX(L,1)
100
105
C
140
200
210
250
260
C
220
C
360
370
380
400
450
 460
 500
550
                                                     DO 710 I=1,N

L=N+1-I

IF(INDEX(L,1)-INDEX(L,2))630,710,630

JRUW=INDEX(L,1)

JCDLUM=INDEX(L,2)

DO 705 K=1,N

SWAP=A(K,JROW)

A(K,JROW)=A(K,JCDLUM)

A(K,JCDLUM)=SWAP

CONTINUE

CONTINUE

CONTINUE

DO 11 K=1,N

IF(IPIVOT(K).NE.1)GU TO 12

CONTINUE

RETURN

WRITE(22,991)
  630
  705
  710
   11
                                                        WRITE(22,991)
FORMAT(/30X, MATRIX IS SINGULAR*/)
   12
991
    740
                                                         RETUKN
                                                         END
   C
```